Primary Care Paramedic

Medical Directives

ALS PCS 5.3



2024 - v3 PRINT DATE 2024-05-015

Introduction

Airway/ Breathing

Cardiac/ Circulation

Level of Consciousness/ Pain/Nausea

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Destination Guidelines The Emergency Health Services Branch of the Ministry of Health Version 5.3 of the ALS Patient Care Standards will now be the standard of care. These standards and guidelines include significant advances to the paramedic scope of practice since they were last published. As the ALS PCS is a living document, this Medical Directive book may not be an accurate reflection of the current scope of practice and/or ALS PCS. Paramedics are to refer to the CPER website for access to the most up to date version of the ALS PCS and to their certification letter for currently authorized medications and procedures.

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Note: This Paramedic guide contains content from the Ministry of Health Advanced Life Support Patient Care Standards, version 5.3 dated February 9, 2024. To access the full document please refer to www.CPER.ca.

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Introduction

PRIMARY CARE PARAMEDIC MEDICAL DIRECTIVES



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Introduction

ADVANCED LIFE SUPPORT PATIENT CARE STANDARDS

Levels of Paramedics

In Ontario, there are 3 levels of qualification for paramedics which lead to Certification as a: Primary Care Paramedic (PCP), Advanced Care Paramedic (ACP), and Critical Care Paramedic (CCP). The qualification for each are set out in Ontario Regulation 257/00 made under the *Ambulance Act*, RSO 1990, c A-19. The qualifications for each include a requirement that the paramedic be authorized by a Medical Director of a Regional Base Hospital (RBH) to perform the controlled acts set out in Schedules 1, 2 and 3 to O. Reg 257/00.

A paramedic may be authorized by the Medical Director to perform controlled acts from the Schedule immediately above their Certification. In this circumstance, the paramedic is required to perform the controlled act to a specific standard as set out in the Advanced Life Support Patient Care Standards (ALS PCS). All advanced medical procedures that are not listed as controlled acts in Schedules 1, 2 and 3, shall also be performed as set out in the ALS PCS.

Purpose of Standards

The ALS PCS reflects current practices for paramedics in Ontario and provides benchmarks for paramedic performance. It also communicates the standards of practice and care by paramedics in Ontario to paramedics, patients, other disciplines and the public in general. In the provision of ALS PCS care, paramedics are required to ensure patient care and documentation is provided in accordance with all appropriate Standards as indicated in O. Reg. 257/00.

Comprehensive Care

Although two patient care standards exist, both Standards represent a continuum of care that is to be followed in an integrated fashion during a call for service. While initiating and continuing treatment prescribed by these Medical Directives, a paramedic must ensure that the patient simultaneously receives care in accordance with the BLS PCS. It is acknowledged that there may be circumstances and situations where complying with ALS PCS is not clinically justified, possible, or prudent (e.g. multiple crews on scene, trapped patient, extenuating circumstances, competing patient care priorities). When treatment deviates from the standards, a paramedic must document the care provided, including reasoning for deviating from the ALS PCS.

Format of the ALS PCS

This document is comprised of a Preamble section and six (6) sections: Section 1 – PCP Core Medical Directives; Section 2 – ACP Core Medical Directives; Section 3 – PCP Auxiliary Medical Directives; Section 4 – ACP Auxiliary Medical Directives; Section 5 – Certification Standard, and Section 6 – Research Trial Standard

Use of the Medical Directives by Paramedics

These Medical Directives apply to paramedics who are authorized by a RBHP Medical Director to provide patient care. Delegation of controlled acts in the ALS PCS to paramedics falls under the exclusive oversight of the RBHP. Critical Care Paramedics and Advanced/Primary Care Flight Paramedics will perform controlled acts in accordance with the Base Hospital (RBHP) Medical Directives issued by the Ornge Base Hospital Medical Director(s).

General Structure of a Medical Directive

All Medical Directives follow the same format and are comprised of the following sections:

Indication: The general medical complaint or problem to which

the Medical Directive applies.

Conditions: Clinical parameters that must be present for a

procedure to be performed or for a medication to be

administered.

Contraindications: Clinical parameters that if present, preclude the

performance of a procedure or the administration of a

medication.

Treatment: Description of the type of procedure to be performed

or the dosing of a medication.

Clinical

Considerations: Key clinical points that provide general guidance to

the proper performance of a procedure or the

administration of a medication.

All of these sections must be taken into account before and during the implementation of a Medical Directive.

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

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Research/ Sp. Proj

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Auxiliary Medical Directives

Additional ("Auxiliary") controlled medical acts or advanced medical interventions may be delegated through use of the Auxiliary Medical Directives. Delegation of Auxiliary Medical Directives by a RBHP Medical Director to paramedics is optional and may be introduced after consultation and mutual agreement between the RBHP and the certified ambulance service that employs the paramedic. Some PCP and ACP Medical Directives contain the phrase, "(if available and authorized)". This phrase qualifies the skill or procedure as optional (i.e. auxiliary) even if included in PCP or ACP Medical Directives

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

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Special Event Medical Directives

Medical Directives have been developed for time limited periods when a mass gathering could potentially strain the resources of the host community. These medical directives shall only be used by paramedics who have completed the necessary training and received Regional Base Hospital Program authorization.

Consent to Treatment in Non-Emergency Situations

Except in emergency circumstances described below, paramedics shall obtain consent prior to administering treatment. If a patient is incapable of consenting to the treatment being proposed by a paramedic, consent may be given or refused on his or her behalf by the patient's substitute decision-maker (SDM). Consent may be expressed or implied. Implied consent may be assumed where a person provides a physical indication that they consent to the treatment being proposed. For example, a patient who cannot speak but extends his hand to a paramedic after the paramedic indicates she is going to perform a simple procedure, such as a blood glucose determination, may be giving implied consent to the treatment plan being proposed.

The elements are required for consent to treatment are:

- a) consent must be given by a person who is capable of giving consent with respect to treatment;
- b) consent must relate to the treatment plan;
- c) consent must be informed;
- d) consent must be given voluntarily; and
- e) consent must not be obtained through misrepresentation or fraud.

Consent to treatment is informed if, before it is given to the person, he or she has:

 a) received the following information that a reasonable person in the same circumstances would require in order to make a decision about the treatment plan;

- i. the nature of the treatment;
- ii. the expected benefits of the treatment;
- iii. the material risks of the treatment:
- iv. the material side effects of the treatment:
- v. alternative courses of action;
- vi. the likely consequences of not having the treatment; and
- b) received responses to his or her requests for additional information about those matters

Valid consent requires that a person has the capacity to provide consent. A person is presumed to have the capacity to provide consent with respect to treatment and a paramedic may rely on that presumption unless the paramedic has reasonable grounds to believe that the person is incapable with respect to the treatment plan. A paramedic must perform a capacity assessment if it is not reasonable in the circumstances to presume the person is capable of consenting to the treatment.

A patient is capable with respect to the treatment plan if the patient is:

- Able to understand the information that is relevant to making a decision about the treatment or alternatives being proposed; and
- Able to appreciate the reasonably foreseeable consequences of a decision or lack of decision with respect to the treatment plan.

If a patient is incapable of consenting to a proposed treatment plan, and the paramedic is aware or is made aware that the person has a prior capable wish with respect to the proposed treatment, they must respect that wish (for example, if the person does not wish to be resuscitated).

Airway /

Breath.

Cardiac/

Circula.

IOC/

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Consent to Treatment in Emergency Situations

Where the person for whom the treatment is being proposed is apparently experiencing severe suffering or is at risk of sustaining serious bodily harm if the treatment is not administered promptly, it is considered to be an emergency.

For situations involving consent to treatment in emergency situations, a paramedic shall comply with the applicable directions contained in the *Basic Life Support Patient Care Standards* (BLS PCS).

Discharge from Care

If a paramedic is certified and authorized by their Regional Base Hospital to perform a prehospital discharge from care as per the applicable Medical Directives, the following applies. For the purpose of the applicable Medical Directives, a patient or substitute decision maker (SDM) present at the scene, must be capable to make an informed decision about their treatment plan.

A paramedic authorized to perform a prehospital discharge from care shall:

- Determine whether a patient may be treated in accordance with the Treat and Discharge component of the applicable Medical Directive,
- Communicate a clinically reasonable differential diagnosis to the patient or SDM,
- 3. Discuss the following elements of a discharge treatment plan:
- a. The clinical situation related to the most likely diagnosis and/or differential diagnoses,
 - b. The symptoms and signs alerting them to seek further medical care (i.e. clues that the condition is worsening or that the diagnosis may not be correct),
 - c. Instructions regarding modifications(s) of activities of daily living following the health event,
 - d. Where possible, provide additional contacts for follow up care,
 - e. Instructions to call 911 back if their condition worsens or recurs, and
- 4. Ensure the patient has the necessary support to follow a discharge treatment plan. These supports may include: a. access to food,
 - b. access to transportation,
 - c. access to alternate health care follow up,
 - d. a safe place to stay,
 - e, responsible adult at the scene available to monitor the patient, and
 - f. consideration of other apparent patient vulnerabilities.

Contact

Refusal of Treatment

If a patient refuses treatment, either in whole or in part, a paramedic shall comply with the applicable directions contained in the BLS PCS.

Airway / Breath.

Intravenous (IV) Access and Therapy by Primary Care Paramedics

There are 2 types of authorization for PCPs IV cannulation and therapy.

"PCP Assist IV" is authorization for a PCP to cannulate a peripheral IV at the request and under the direct supervision of an ACP. The patient must require a peripheral IV in accordance with the indications listed in the Intravenous and Fluid Therapy Medical Directive - Auxiliary. The ACP will perform all IV therapy in accordance with the Intravenous and Fluid Administration Medical Directive once intravenous access is obtained. PCPs authorized in PCP Assist IV are not authorized to administer IV therapy.

This authorization level can no longer be obtained and only those who have previously received this authorization may continue to practice at this level.

"PCP Autonomous IV" is authorization for a PCP to independently cannulate an IV according to the Intravenous and Fluid Therapy Medical Directive – Auxiliary. PCPs authorized in PCP Autonomous IV are authorized to administer IV therapy according to applicable Medical Directives.

Authorization for each type shall meet the requirements established by the OBHG MAC.

Home Medical Technology and Novel Medications

As community care advances, new home medical technologies and novel medications are being introduced for home use by patients and caregivers trained in the care required. They are generally used by patients with complex medical histories who may require emergent interventions which are not described in, or aligned with, the BLS PCS or ALS PCS.

A "home medical technology" is an external or internal mechanical device prescribed by a member of a regulated health profession for the purpose of treating a medical condition.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic.

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Intro

Airway / Breath. A "novel medication" is a self/caregiver-administered medication prescribed by a member of a regulated health profession that is required to treat patients with generally rare and unusually complex chronic medical conditions which are often end stage. The medication may be self/caregiver-administered by any route into any part of the body.

Cardiac/

A paramedic may accept the claim that a patient or caregiver has knowledge and training about the technology or medication encountered. A paramedic may only assist a patient or caregiver within the authorized paramedic skill set.

LOC/ Pain/ Nausea

Proced

For unusual circumstances requiring interventions in the out of hospital setting, the RBH may create local training modules, treatment guidelines or medical directives

Patching

A paramedic shall patch to the Base Hospital when:

- a) a medical directive contains a mandatory provincial patch point; OR
- b) for situations that fall outside of these Medical Directives where the paramedic believes the patient may benefit from online medical direction that falls within the prescribed paramedic scope of practice; OR
- for consultation when, in the paramedics opinion the patient presentation or situation warrants and medical advice is required.

Research/ Sp. Proj

Medical

Refer.

In cases where a treatment option requires the prior authorization by the BHP AND the BHP cannot be reached despite reasonable attempts by the paramedic to establish contact, a paramedic may initiate the required treatment without the requisite online authorization if the patient requires a critical, potentially life-saving, intervention and, in the paramedic's opinion, the intervention would otherwise apply. All patch failures must be reported in a timely manner to the RBHP in accordance with local policy and procedures. Paramedics should document the attempts to patch to the BHP on the Ambulance Call Report (ACR).

Medic. Info.

If a BHP directs a paramedic to perform an assessment or intervention that exceeds the paramedic's scope of practice, the paramedic must advise the BHP of such and notify the physician that they cannot comply with the direction as it exceeds their scope of practice. In such cases, a paramedic should ask the BHP to provide alternative direction.

Contact

Incident Reporting

Paramedics shall adhere to their ambulance service policies and the Ontario Ambulance Documentation Standards (incorporated by reference in Ontario Regulation 257/00) for incident reporting. Paramedics shall also adhere to additional RBH policies regarding reporting of clinical care incidents to the RBHP.

Airway / Breath.

Responsibility of Care

Each paramedic is equally responsible for patient care within their scope of practice. If the care exceeds a paramedics scope of practice, responsibility for that continued care shifts to the higher certified paramedic.

Cardiac/ Circula.

If there is any disagreement between paramedics, the Base Hospital physician may be contacted. It is expected that when an intervention has been performed, the paramedic most appropriate for that intervention will remain responsible for the patient. LOC/ Pain/ Nausea

The risks to the patient during transport should be assessed and discussed prior to transferring care from a higher to lower level of paramedic (e.g.: ACP to PCP), paramedics must alert the highest-level paramedic of any change of patient status at any time in the call.

Proced.

When transferring care from one level of paramedic to another, paramedics shall provide:

Research/ Sp. Proj

- a) current CTAS level;
- b) a history of the patient's current problem(s) and relevant past medical history;
- c) pertinent physical findings;
- d) a summary of management at scene/en route;
- e) the patient's response to treatment, including most recent vital signs; and
- f) the reason for transfer in cases of inter-facility transfers.

Medical Refer.

The transfer of responsibility of patient care is a critical juncture along the clinical care continuum. When transferring patient care to another health care provider (e.g. nurse, physician, etc.), a paramedic must comply with the BLS PCS regarding such transfers

Medic. Info.

Research

Clinical research is fundamental to the practice of medicine and the development of safer, more effective treatment options for patients. At times, research protocols

Contact

Intro

require temporary changes to patient care standards. Changes to patient care standards will be approved and introduced by the MOH.

Airway / Breath.

Patient Care Model

Cardiac/ Circula. Any patient care model subject to The Patient Care Model Standard (PCMS) requires approvals and training as per the PCMS. Paramedics shall assess and provide treatment to all patients in accordance with the ALS PCS and BLS PCS when patients do not completely meet the specific parameters of approved Patient Care Models.

LOC/ Pain/ Nausea

Conventions

"Conventions" refers to a consistent application of terms throughout the Medical Directives based on definitions below.

Proced.

The word 'consider' is used repeatedly throughout the Medical Directives. Where this word appears, it indicates that a paramedic shall initiate the treatment when the indications are first identified unless there is strong clinical rationale to withhold or delay treatment or other extenuating circumstances. A paramedic must document his or her justification for withholding treatment on the ACR.

Research/ Sp. Proj

Medication Doses and Administration

Unless specified within the medical directive, the number of recommended medication doses may be administered regardless of any previous administrations. When more than one route of medication administration is listed, clinical circumstances for each case should determine the final route chosen.

Medical Refer.

When more than one route of medication administration is listed, the order of preference for route of administration is from left to right. Clinical circumstances for each case should determine the final route chosen.

Medic. Info. Pediatric medication doses can vary slightly according to the source of expert opinion. The pediatric medication doses in the ALS PCS are the preferred doses. However, medication doses as determined by an up-to-date version of a widely accepted RBHP approved pediatric emergency tape (e.g. Broselow Tape) are an acceptable alternative. Use of a pediatric emergency tape shall be documented on the ACR when it is used to determine a pediatric medication dose.

Contact

Medication doses may be calculated based upon weight or other factors and result in a fraction that cannot be measured accurately. In these cases, the medication dose delivered will be rounded to the closest dose that can accurately be measured

Age and Vital Signs

The general age cut off between adults and pediatrics is 18 years (under 18 yrs. is generally considered a pediatric patient). There is a wide range of "normal" for vital signs in adults and especially pediatrics. As much as possible, ages for pediatrics and cut off points for vital signs have been kept consistent throughout the Medical Directives. However, clinical research and expert opinion have resulted in a number of exceptions which in each case has been deliberately chosen and is clearly noted in each Medical Directive. Age will be written as a number of hours, days, or years throughout the medical directives. There is a deliberate gap in the definition of normotension and hypotension in adults.

ADULTS

Normotension SBP ≥100mmHg

Hypotension SBP <90 mmHg

Heart rate Heart rate is always in beats per minute according to a cardiac monitor when it is applied. In situations where a cardiac monitor is not indicated then the heart rate is equal to the pulse rate.

Bradycardia HR <50 BPM

Tachycardia HR ≥100 BPM

Tachypnea RR ≥28 breath/min

PEDIATRICS

Age	Respiratory Rate	Heart Rate
0-3 months	30-60	90-180
3-6 months	30-60	80-160
6-12 months	25-45	80-140
1-3 yr	20-30	75-130
6 yr	16-24	70-110
10 yr	14-20	60-90

Normotension SBP \geq 90 mmHg + (2 x age in years) **Hypotension** SBP < 70 mmHg + (2 x age in years) Airway / Breath.

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Weight (kg) (age x 2) + 10

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Medic. Info.

Contact

HYPOGLYCEMIA

Age	Blood glucose level
<2 yr	<3.0 mmol/L
≥2 yr	<4.0 mmol/L

Level of Awareness (LOA):

The word 'altered' refers to a GCS that is less than normal for the patient.

The word 'unaltered' refers to a GCS that is normal for the patient. This may be a GCS <15.

Commonly Used Abbreviations

The following abbreviations, in alphabetical order, appear in the Advanced Life Support Patient Care Standards:

Α

ACP	Advanced Care Paramedic
ALS	Advanced Life Support

ALS PCS Advanced Life Support Patient Care Standards

ASA Acetylsalicylic acid

AED automated external defibrillation

В

BHP Base Hospital Physician

BLS PCS Basic Life Support Patient Care Standards

BPM Beats per minute BVM Bag-valve-mask

C

CCP Critical Care Paramedic

COPD Chronic obstructive pulmonary disease

COWS Clinical Opiate Withdrawal Scale

cm Centimeter

CPAP Continuous positive airway pressure

Airway / Breath.

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Medical Refer.

Medic. Info.

CPR CTAS CVA CVAD	Cardiopulmonary Resuscitation Canadian Triage and Acuity Scale Cerebral vascular accident Central venous access device
D	
DKA DNR	Diabetic ketoacidosis Do Not Resuscitate
E	
ECG ED ETCO ₂ ETT	Electrocardiogram Emergency Department End tidal carbon dioxide Endotracheal tube
F	
FiO ₂	Fraction of inspired oxygen
G	
g GCS gtts	Gram Glasgow Coma Scale Drops
Н	
H ₂ O HR Hx HF	Water Heart rate History Hydrofluoric Acid
IM	Intramuscular
IN IO	Intranasal Intraosseous
IV	Intravenous
J	
j	Joule
K	
kg	Kilogram

Destinat. Guide.

Intro	L	
Airway / Breath.	LOA LOC M Max.	Level of awareness Level of consciousness Maximum
Cardiac/ Circula.	MAC mcg MDI mg Min.	Medical Advisory Committee Microgram Metered dose inhaler Milligram Minimum
LOC/ Pain/ Nausea	min mL/kg mmHg MOH ms	Minute Milliliter per kilogram Millimeters of mercury Ministry of Health Milliseconds
Proced.	N/A NaCl nare	Not applicable Sodium chloride Nostril
Research/ Sp. Proj	NEB NPA NSAID	Nebulized Nasopharyngeal airway Non-steroidal anti-inflammatory drug
Medical Refer.	OBHG-MAC	Ontario Base Hospital Group - Medical Advisory Committee Oropharyngeal airway
Medic. Info.	PCP PEA PPV PO	Primary Care Paramedic Pulseless electrical activity Positive Pressure Ventilation by mouth/oral
Contact	PRN Q q	as needed every

R		iiido
RBHP ROSC RR	Regional Base Hospital Program Return of spontaneous circulation Respiratory rate	Airway / Breath.
SAED SC SL SBP SpO ₂	Semi-automatic external defibrillation Subcutaneous Sublingual Systolic blood pressure Saturation of peripheral oxygen	Cardiac/ Circula.
STEMI T TBI	ST-segment elevation myocardial infarction Traumatic brain injury	LOC/ Pain/ Nausea
TCP TOP TOR	Transcutaneous pacing Topical Termination of Resuscitation	Proced.
V VF	Upper respiratory tract infection Ventricular Fibrillation	Research/ Sp. Proj
VF VT VSA W	Ventricular Florination Ventricular Tachycardia Vital signs absent	Medical Refer.
WNL	Within normal limits	Medic.

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Reference and Educational Notes

The RBHPs have created a companion document of reference and educational notes intended to assist paramedics in implementing these Medical Directives. This will facilitate regular updating of these notes without having to issue frequent changes to the standards. It is expected that paramedics have mastered the relevant information as part of initial training and certification and have maintained their knowledge through continuing education and self assessment and reflective practice. The reference and educational notes do not define a standard of care and is not a nested document to this standard; however, they should be considered useful in ensuring that an appropriate standard of care is met.

Airway/Breathing

PRIMARY CARE PARAMEDIC MEDICAL DIRECTIVES



Breath.

Bronchoconstriction Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized

INDICATIONS

Cardiac/ Circula.

IOC/

Pain/ Nausea

Proced

Respiratory distress:

AND

Suspected bronchoconstriction

CONDITIONS

Salbutamol

AGE: N/A LOA: N/A

HR: N/A RR· N/A

SBP: N/A

Research / Other: N/A Sp. Proj

EPINEPHrine

AGE: N/A WEIGHT: N/A LOA: N/A N/A HR.

RR. **BVM** ventilation required

SBP: N/A

Other: Hx of asthma

Dexamethasone

AGE: N/A LOA: N/A HR: N/A RR. N/A SBP: N/A

Other: Hx of asthma OR

COPD OR

20 pack-year history

of smoking

Medical Refer.

Medic. Info

CONTRAINDICATIONS Salbutamol

Allergy or sensitivity to salbutamol

EPINEPHrine

Allergy or sensitivity to **FPINFPHrine**

Dexamethasone

Allergy or sensitivity to steroids

Currently on PO or parenteral steroids

Contact

TREATMENT



Patient Drug Dose Route Time.

Consider salbutamol

	We	ight	Weight			
	<25	i kg	≥25 kg			
	Route	Route	Route	Route		
	MDI* NEB		MDI*	NEB		
Dose	Up to 600 mcg (6 puffs)	2.5 mg	Up to 800 mcg (8 puffs)	5 mg		
Max. Single Dose	600 mcg	2.5 mg	800 mcg	5 mg		
Dosing interval	5-15 min. PRN	5-15 min. PRN	5-15 min. PRN	5-15 min. PRN		
Max. # of doses	3	3	3	3		

^{* 1} puff=100mcg

Consider EPINEPHrine

IM Concentration 1 mg/mL = 1:1,000 0.01 mg/kg**

Route

Dos	se	0.01 mg/kg**
Max. sing	gle dose	0.5 mg
Dosing in	nterval	N/A
Max. # 0	f doses	1

^{**} The EPINEPHrine dose may be rounded to the nearest 0.05 mg.

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Consider dexamethasone			
		Route	
		PO/IM/IV	
	Dose	0.5 mg/kg	
	Max. single dose	8 mg	
	Dosing interval	N/A	
	Max. # of doses	1	

CLINICAL CONSIDERATIONS

EPINEPHrine should be the 1st medication administered if the patient is apneic. Salbutamol MDI may be administered subsequently using a BVM MDI adapter.

Nebulization is contraindicated in patients with a known or suspected fever or in the setting of a declared febrile respiratory illness outbreak by the local medical officer of health.

When administering salbutamol MDI, the rate of administration should be 100 mcg approximately every 4 breaths.

A spacer should be used when administering salbutamol MDI.

EPINEPHrine 1 mg/mL = 1:1000 IM Dosing Chart

Dose (0.01 mg/kg) is rounded to the nearest 0.05mg

Use a 1 mL syringe							
AG	iE V	VEI	GHT	DOSE	(mg)	VOLUME (mL) (rounded)	
3 m	onths	5	kg	0.05	mg	0.05	mL
6 m	onths	8	kg	0.08	mg	0.10	mL
9 m	onths	10	kg	0.10	mg	0.10	mL
1 ye	ear	12	kg	0.12	mg	0.10	mL
2 ye	ears	14	kg	0.14	mg	0.15	mL
3 ye	ears	16	kg	0.16	mg	0.15	mL
4 ye	ears	18	kg	0.18	mg	0.20	mL
5 ye	ears	20	kg	0.20	mg	0.20	mL
6 ye	ears :	22	kg	0.22	mg	0.20	mL
7 ye	ears	24	kg	0.24	mg	0.25	mL
8 ye	ears	26	kg	0.26	mg	0.25	mL
9 ye	ears	28	kg	0.28	mg	0.30	mL
10 ye	ears	30	kg	0.30	mg	0.30	mL
11 ye	ears	32	kg	0.32	mg	0.30	mL
12 ye	ears	34	kg	0.34	mg	0.35	mL
13 ye	ears	36	kg	0.36	mg	0.35	mL
14 ye	ears	38	kg	0.38	mg	0.40	mL
Adı	ult	50	kg	0.50	mg	0.50	mL

Note: Dosage administered can be calculated by the weight based calculation in the Medical Directive and/or by using the above chart. Administered dosage in the chart may be rounded to the nearest volume increment that can be accurately measured. Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer.

Medic.

Contact

Destinat.

Breath.

Cardiac/ Circula

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

Medic Info.

Contact

Moderate to Severe Allergic Reaction **Medical Directive**

A Primary Care Paramedic may provide the treatment prescribed in this medical directive if authorized

INDICATIONS

Exposure to a probable allergen;

AND

Signs and/or symptoms of a moderate to severe allergic reaction (including anaphylaxis)

CONDITIONS

EPINEPHrine

AGE: N/A WFIGHT: N/A

I OA· N/A HR· N/A

RR· N/A SBP: N/A

Other: For anaphylaxis

only

DiphenhydrAMINE

AGE: N/A

WEIGHT: ≥25 kg

I OA· N/A HR: N/A RR: N/A

SBP: N/A Other: N/A

CONTRAINDICATIONS

EPINEPHrine

Alleray or sensitivity to **FPINFPHrine**

DiphenhydraMINE

Alleray or sensitivity to diphenhydraMINE

TREATMENT



Patient Drug Dose Route Time.

Consider EPINEPHrine

	Route
	IM
	Concentration
	1 mg/mL = 1:1,000
	0.01 mg/kg*
se	0.5 mg

Dose	0.01 mg/kg*
Max. single dose	0.5 mg
Dosing interval	Minimum 5 min
Max. # of doses	2

*The EPINEPHrine dose may be rounded to the nearest 0.05 mg.

Consider diphenhydraMINE:

	Weight	Weight
	≥25 kg to <50 kg	≥50 kg
	Route	Route
	IV/IM	IV/IM
Dose	25 mg	50 mg
Max. single dose	25 mg	50 mg
Dosing interval	N/A	N/A
Max. # of doses	1	1

Airway /

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

CLINICAL CONSIDERATIONS

Airway / Breath. EPINEPHrine administration takes priority over IV access.

 $\ensuremath{\mathsf{IV}}$ administration of diphenhydra MINE applies only to PCPs authorized for PCP Autonomous IV.

Cardiac/ Circula. NOTE: Refer to page 24 for EPINEPHrine 1 mg/mL = 1:1000 IM Dosing Chart.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

Medic.

Contact

Croup Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized.

INDICATIONS

Current history of URTI;

AND

Barking cough or recent history of a barking cough

CONDITIONS

EPINEPHrine

AGE: ≥ 6 months to <8 years

LOA: N/A

HR: <200 bpm RR: N/A SBP: N/A

Other: Stridor at rest

Dexamethasone

AGE: ≥ 6 months to <8 years

LOA: Unaltered HR: N/A RR· N/A SBP: N/A

Other: For mild, moderate and severe croup

CONTRAINDICATIONS

EPINEPHrine

Allergy or sensitivity to EPINEPHrine

Dexamethasone

Allergy or sensitivity to steroids Steroids received within the last 48 hours Unable to tolerate oral medications

Contact

Breath.

Cardiac / Circula.

LOC/ Pain/ Nausea

Proced

Research / Sp. Proi

Medical Refer.

Medic. Info

Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

TREATMENT

Patient Drug Dose Route Time.

Consider EPINEPHrine

	Weight	Weight
	<10 kg	≥10 kg
	Route	Route
	NEB	NEB
	Concentration	Concentration
	1 mg/mL = 1:1,000	1 mg/mL = 1:1,000
Dose	2.5 mg	5 mg
Max. single dose	2.5 mg	5 mg
Dosing interval	N/A	N/A
Max. # of doses	1	1

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Consider dexamethasone

	Age ≥ 6 months to < 8 years
	Route
	PO
Dose	0.5 mg/kg
Max. single dose	8 mg
Dosing interval	N/A
Max. # of doses	1

CLINICAL CONSIDERATIONS

N/A

Croup Assessment

- Croup is an upper respiratory infection that is generally the result of a viral infection
- It tends to occur in children aged 6 months to 3 years, and is most prevalent at the age of 2 years.
- It is characterized by swelling and irritation of the respiratory tract, and is often associated with a "barking style" cough.
- The severity of the symptoms can be characterized using the guideline below.
- Generally speaking, patients with moderate to severe croup should be considered for therapy as per the Medical Directive.

WESTLEY CROUP SCORE:

This allows the severity of symptoms to be classified. Maximum score possible is 17.

		Score				
	0	1	2	3	4	5
Inspiratory Stridor	-	Audible with stethoscope	Audible without stethoscope	-	-	-
Retraction	-	Mild	Moderate	Severe	-	-
Air entry	Normal	Decreased	Severely decreased	-	-	-
Cyanosis	None	-	-	-	With agitation	At rest
Conscious level	Normal	-	-	-	-	Altered

Score of 2-3: Indicates mild croup.

Score of 4-7: Indicates moderate croup.

Score of >7: Indicates severe croup.

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic.

Intro

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

Medic.

Contact

Continuous Positive Airway Pressure (CPAP) Medical Directive - AUXILIARY

A Primary Care Paramedic may provide the treatment prescribed in this auxiliary Medical Directive if authorized.

INDICATIONS

Severe respiratory distress;

AND

Signs and/or symptoms of acute pulmonary edema or COPD.

CONDITIONS

CPAP

AGE: ≥18 years LOA: N/A HR: N/A RR: Tachypnea

Other: SpO₂ < 90% or accessory muscle use

CONTRAINDICATIONS

SBP: Normotension

CPAP

Asthma exacerbation

Suspected pneumothorax

Unprotected or unstable airway

Major trauma or burns to the head or torso

Tracheostomy

Inability to sit upright

Unable to cooperate

TREATMENT

Initial Setting 5 cm H	Or equivalent flow 2O rate of device as per BH direction
Titration increment 2.5 cm l	Or equivalent flow rate of device as per BH direction
Titration interval 5 min	1
Max. setting 15 cm H	Or equivalent flow rate of device as per BH direction

Initial FiO ₂	50-100%
FiO₂ increment	SpO ₂ <92% despite treatment and/or
(if available on device)	10 cm H ₂ O pressure or equivalent flow rate of
1.7	device as per BH direction
Max FiO ₂	100%

Confirm CPAP pressure by manometer (if available)

CLINICAL CONSIDERATIONS

N/A

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Intro

Airway / Breath.

Cardiac/

LOC/ Pain/ Nausea

Proced.

Research/ Sp.Proj

Medical Refer.

Medic.

Contact

Destinat. Guide.

Supraglottic Airway Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized.

INDICATIONS

Need for ventilatory assistance or airway control;

AND

Other airway management is ineffective.

CONDITIONS

Supraglottic Airway

AGE: N/A LOA: N/A HR: N/A RR: N/A SBP: N/A

Other: Absent gag reflex

CONTRAINDICATIONS

Supraglottic Airway

Airway obstructed by a foreign object Known esophageal disease (varices)

Trauma to the oropharynx

Caustic ingestion

TREATMENT

Consider supraglottic airway insertion

The maximum number of supraglottic airway insertion attempts is 2.

Confirm supraglottic airway placemen	t
Method	Method
Primary	Secondary
ETCO ₂ (Waveform capnography)	ETCO ₂ (Non-waveform device)
	Auscultation
	Chest rise
	Auscultation

CLINICAL CONSIDERATIONS

An attempt at supraglottic airway insertion is defined as the insertion of the supraglottic airway into the mouth.

Confirmation of supraglottic airway should use ETCO₂ (Waveform capnography). If waveform capnography is not available or is not working, then at least 2 secondary methods must be used.

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic.

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer.

Medic. Info.

Contact

Endotracheal and Tracheostomy Suctioning & Reinsertion Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized.

INDICATIONS

Patient with endotracheal or tracheostomy tube;

AND

Airway obstruction or increased secretions

CONDITIONS

Suctioning

AGE: N/A LOA: N/A HR: N/A

RR: N/A SBP: N/A

Other: N/A

Emergency Tracheostomy Reinsertion

AGE: N/A
LOA: N/A
HR: N/A
RR: N/A
SBP: N/A

Other: Patient with an existing tracheostomy where the inner and/or outer cannula(s) have been removed from the airway **AND**

Respiratory distress AND

Inability to adequately ventilate **AND** Paramedics are presented with a tracheostomy cannula for the identified patient

CONTRAINDICATIONS

Suctioning

N/A

Emergency Tracheostomy reinsertion

Inability to landmark or visualize

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TREATMENT

	< 1 year	≥ 1 year to < 12 years	≥ 12 years
Dose	Suction at 60-100 mmHg	Suction at 100-120 mmHg	Suction at 100-150 mmHg
Max. single dose	10 seconds	10 seconds	10 seconds
Dosing interval	1 minute	1 minute	1 minute
Max. # of doses	N/A	N/A	N/A

Consider emergency tracheostomy reinsertion

The maximum number of attempts is 2.

CLINICAL CONSIDERATIONS

Suctioning:

Pre-oxygenate with 100% oxygen.

In an alert patient, whenever possible, have patient cough to clear airway prior to suctioning.

Emergency tracheostomy reinsertion:

A reinsertion attempt is defined as the insertion of the cannula into the tracheostomy. A new replacement inner or outer cannula is preferred over cleaning and reusing an existing one.

Utilize a family member or caregiver who is available and knowledgeable to replace the tracheostomy cannula.

Breath.

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced

Research/ Sp. Proj

Medical Refer.

Medic. Info

Contact

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Destinat. Guide.

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer.

Medic. Info.

Contact

Destinat. Guide.

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Destinat. Guide.

Cardiac/Circulation

PRIMARY CARE PARAMEDIC MEDICAL DIRECTIVES



Airway / Breath.

Cardiac/

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Destinat.

Medical Cardiac Arrest Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized.

INDICATIONS

Non-traumatic cardiac arrest.

PRIMARY CLINICAL CONSIDERATION(S)

In the following settings, consider very early transport after a minimum of one analysis (and defibrillation if indicated) once an egress plan is organized:

- pregnancy presumed to be ≥ 20 weeks gestation (fundus above umbilicus, ensure manual displacement of uterus to left):
- 2. hypothermia;
- 3. airway obstruction;
- 4. non-opioid drug overdose/toxicology, or;
- 5. other known reversible cause of arrest not addressed.

For patients in refractory VF or pulseless VT, transport of the patient should begin after the third consecutive shock. Refractory VF or pulseless VT is defined for the purpose of this directive, as persistent VF or pulseless VT after 3 consecutive shocks.

CONDITIONS

CPR

AGE: N/A LOA: Altered HR: N/A

SBP: N/A

Other: Performed in 2

Manual Defibrillation

AGE: ≥ 24 hours
LOA: Altered
HR: N/A
RR: N/A

SBP: N/A
Other: VF **OR** pulseless

VT

AED or SAED

AGE: ≥ 24 hours

HR: N/A RR: N/A SBP: N/A

Other: Defibrillation indicated

If not using manual defibrillation

EPINEPHrine

AGE: ≥ 24 hours LOA: Altered HR: N/A RR: N/A SBP: N/A

Other: Anaphylaxis suspected as causative event

Medical TOR

AGE: ≥ 16 years LOA: Altered HR: N/A RR: N/A SBP: N/A

Other: Arrest not witnessed by paramedic **AND** No ROSC after 20 minutes of resuscitation **AND**

No defibrillation delivered

CONTRAINDICATIONS

CPR

Obviously dead as per

Meet conditions of the BLS PCS Do Not Resuscitate (DNR) Standard Manual Defibrillation

Rhythms other than VF or pulseless VT

AED or SAED Defibrillation

Non-shockable rhythm

Airway / Breath

Intro

Cardiac/

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

Medic.

Contact

Airway / Breath.

EPINEPHrine

Allergy or sensitivity to EPINEPHrine

Medical TOR

Age

Known reversible cause of the arrest unable to be addressed

Pregnancy presumed to be ≥ 20 weeks gestation

Suspected hypothermia

Airway obstruction

Non-opioid drug overdose/toxicology

Cardiac/ Circula.

TREATMENT



Patient Drug Dose Route Time.

Proced.

Consider CPR as per current Heart and Stroke Foundation of Canada Guidelines

Age

Research/ Sp. Proj Consider **Manual defibrillation** (if available and authorized)

>24 hours to ≥8 years <8 years Dose 1 defibrillation 1 defibrillation First dose As per RBHP / manufacturer 2 J/kg Subsequent and 4 J/kg As per RBHP / manufacturer max. dose(s) Dosing interval 2 min 2 min Max. # of doses N/A N/A

Medical Refer.

Medic. Info.

Contact

Destinat.

Consider AED or SAED defibrillation (if not using manual defibrillation)

Age	Age
≥24 hours to <8 years	≥8 years
1 defibrillation with or without pediatric attenuator cable	1 defibrillation
As per RBHP / manufacturer	As per RBHP / manufacturer
2 min	2 min
N/A	N/A
	1 defibrillation with or without pediatric attenuator cable As per RBHP / manufacturer 2 min

Consider **EPINEPHrine** (only if anaphylaxis is suspected as causative event)

	Route
	IM
	Concentration
	1 mg/mL = 1:1,000
Dose	0.01 mg/kg*
Max. single dose	0.5 mg
Dosing interval	N/A
Max. # of doses	1

^{*} The EPINEPHrine dose may be rounded to the nearest 0.05 mg

Δ Mandatory Provincial Patch Point Δ

Patch to consider Medical TOR (if applicable).

If the patch fails or if Medical TOR does not apply, transport to the closest appropriate hospital following ROSC or 20 minutes of resuscitation without ROSC.

Patch early (e.g. following the 4th analysis) to consider TOR if there are extenuating circumstances surrounding egress, prolonged transport or significant clinical limitations where the paramedic considers ongoing resuscitation to be fulfile

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer

Medic.

Contact

Airway / Breath

Cardiac/ Circula

IOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer

Medic Info.

Contact

Destinat

CLINICAL CONSIDERATIONS

Consider regional base hospital advanced airway strategy (e.g. SGA medical directive) where more than OPA/NPA and BVM is required.

There is no clear role for routine administration of naloxone in confirmed cardiac arrest

The BHP might not authorize TOR even though the patient meets TOR rule. Factors may include: location of the patients, EtCO2, age, bystander witnessed, bystander CPR, transportation time, and unusual cause of cardiac arrest such as electrocution, hanging, and toxicology.

The BHP may authorize TOR even though the patient does **not** meet the TOR rule. Factors that may be taken into account include extenuating egress limitations. prolonged transport, caregiver wishes, existence of DNR confirmation form, and underlying end stage progressive illness.

LOCAL BHP CONSULTATION ADVISORY

In extenuating circumstances during unusual or prolonged codes, Paramedics may choose to patch for consultation.

Extenuating circumstances may include, but are not limited to, the following:

- 1. Unusual cardiac arrest causes (ie. FBAO, hypothermia, electrocution, toxicity)
- 2. Excessive epinephrine administration (>5-6mg) in prolonged resuscitations.
- 3. Excessive number of shocks (>3 with vector change) delivered without change in refractory dysrhythmia.

Patient presentation/underlying cause of cardiac arrest should be considered when carrying out a treatment plan.

NOTE: Refer to page 46 for Defibrillation Joule Setting Reference Chart.

NOTE: Refer to page 120 for CPR Guidelines

Pediatric Defibrillation Joule Setting Chart

Age	Approx Weight	First Defib Setting (2J/kg)	Subsequent Defib Setting (4J/kg)
0 to 30 days		N/A	N/A
≥1 month to <3 months	<5kg	10 J	20 J
≥3 months to <1 year	≥5 to <12kg	15 J	30 J
≥1 to <5 years	≥12 to <20kg	30 J	70 J
≥5 to <8 years	≥20 to <30kg	50 J	100 J
≥8 years		Adult Manual De settings	fibrillation

Adult Defibrillation Joule Settings Reference

Manufacturer:	Series:	Joule Settings:
Medtronic	Lifepack	200, 300, 360 Joules
Phillips	MRX / FR2	150 Joules non escalating
ZOLL	E, M, or X Series	120, 150, 200 Joules

Airway / Breath.

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer.

Medic. Info.

Contact

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp.Proj

Medical Refer.

Medic. Info.

Contact

Trauma Cardiac Arrest Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized.

INDICATIONS

Cardiac arrest secondary to severe blunt or penetrating trauma.

CONDITIONS

CPR

AGE: N/A LOA: Altered HR: N/A RR: N/A SBP: N/A

Other: Performed in 2 minute intervals

AED or SAED Defibrillation

AGE: ≥24 hours
LOA: Altered
HR: N/A
RR: N/A
SBP: N/A
Other: Defibrillation
indicated

Manual Defibrillation

AGE: ≥24 hours LOA: Altered HR: N/A RR: N/A

Other: VF **OR** pulseless VT

Trauma TOR

AGE: ≥16 years LOA: Altered HR: 0

RR: 0 SBP: N/A

Other: No palpable pulses AND

No defibrillation delivered AND

Rhythm Asystole AND

No signs of life at any time since fully extricated OR

Signs of life when fully extricated with the closest ED \geq 30 min transport

time away OR

Rhythm PEA with the closest ED ≥30 min transport time away.

CONTRAINDICATIONS

CPR

Obviously dead as per BLS PCS

Meet conditions of Do Not Resuscitate (DNR) Standard

AED or SAED Defibrillation

Non-shockable rhythm

Manual Defibrillation

Rhythms other than VF or pulseless VT

Trauma TOR

Age <16 years

Defibrillation delivered

Signs of life at any time since fully extricated medical contact

Rhythm PEA and closest ED <30 min transport time away

Patients with penetrating trauma to the torso or head/neck and Lead Trauma Hospital < 30 min transport time away

Intro

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic.

Contact

TREATMENT

Airway / Breath.

Consider CPR as per the current Heart and Stroke Foundation of Canada guidelines

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Consider Manual defibrillation (if available and authorized)

	Age	Age
	≥24 hours to <8 years	≥8 years
Dose	1 defibrillation	1 defibrillation
Initial dose	2 J/kg	As per RBHP / manufacturer
Dosing interval	N/A	N/A
Max. # of doses	1	1

Consider AED or SAED defibrillation (if not using manual defibrillation)

≥24 hours to <8 years	≥8 years
1 defibrillation With or without Pediatric Attenuator Cable	1 defibrillation
As per RBHP / manufacturer	As per RBHP / manufacturer
N/A	N/A
1	1
	1 defibrillation With or without Pediatric Attenuator Cable As per RBHP / manufacturer

Mandatory Provincial Patch Point 🛆

Patch to BHP for authorization to apply the Trauma TOR if applicable. If the BHP patch fails, or the Trauma TOR does not apply, transport to the closest appropriate receiving facility following the 1st analysis/defibrillation.

Meets Cardiac arrest obviously death criteria or secondary to Follow conditions of DNR BLS PCS severe blunt or penetrating standard trauma ·Nο Simultaneously address reversible causes Hypovolemia: control external hemorrhage. CPR (through out duration of call) splint pelvis/fractures Oxygenation: Prioritize basic airway management and ventilation to correct hypoxia, use advanced AW only as needed Apply defib pads to all patients ≥24 hours of age VF / VT Determine Rhythm Asystole (Shockable) Patient with penetrating trauma to the torso, head or PEA Defibrillation x1 neck and lead trauma (no shock) Yes Hospital <30min away Νo Pt.≥16 years of age? Yes Signs of life Yes No Drive time Patch to closest ED <30 min TOR granted? Νo Yes TOR implemented Transport to Emergency Department

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

CLINICAL CONSIDERATIONS

Airway / Breath

Cardiac/

Circula

IOC/ Pain/

Nausea

If no obvious external signs of significant blunt trauma, consider medical cardiac arrest and treat according to the appropriate medical cardiac arrest directive.

Signs of life: specifically any spontaneous movement, respiratory efforts, organized electrical activity on ECG, and reactive pupils.

An intravenous fluid bolus may be considered, where it does not delay transport and should not be prioritized over management of other reversible pathology.

NOTE: Refer to page 46 for Defibrillation Joule Setting Reference Chart.



NOTE: Refer to page 120 for CPR Guidelines

Proced.

Research/ Sp. Proj

Medical Refer

Medic Info.

Contact

Newborn Resuscitation Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized

INDICATIONS

Newborn patient.

CONDITIONS

Positive Pressure Ventilation (PPV)

AGE: <24 hours I OA· N/A

HR: < 100 bpm RR: N/A

SRP: N/A Other: N/A

CPR

AGE: <24 hours LOA: N/A HR· < 60 bpm RR· N/A

SRP: N/A

Other: After 30 seconds of PPV using

room air

CONTRAINDICATIONS

Positive Pressure Ventilation (PPV)

Obviously dead as per BLS PCS

Presumed gestational age less than 20 weeks

CPR

Obviously dead as per BLS PCS

Presumed gestational age less than 20

weeks

Intro

Airway / Breath

Cardiac/ Circula

IOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer

Medic Info.

Contact

TREATMENT

Airway / Breath.

Consider PPV as per the treatment flowchart

Consider CPR as per the current Heart and Stroke Foundation of Canada Guidelines

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

Medic. Info.

Contact

Newborn Patient Targeted Preductal SpO₂ (Right Hand) Provide warmth: 1 min 60% - 65% Term gestation and clear airway as necessary 2 min 65% - 70% good muscle tone and (do not routinely suction) 3 min 70% - 75% breathing or crying? dry; and ongoing evaluation 75% - 80% 4 min 5 min 80% - 85% 10 min 85% - 95% No For 30 seconds: 1. Provide warmth; 2. Position/clear airway; and 3. Dry. stimulate, reposition Evaluate respirations and heart rate (consider use of SpO₂ and cardiac monitoring) Breathing and HR ≥ 100 bpm? Supportive Care Yes For 30 seconds: 1. Provide positive pressure ventilation (BVM) using room air HR ≥ 60 bpm? Yes No For 30 seconds: No 1. Administer chest compressions; and 2. Provide positive pressure ventilation (BVM) using 100% oxygen HR ≥ 60 bpm?

Destinat.

CLINICAL CONSIDERATIONS

If newborn resuscitation is required, initiate cardiac monitoring and right-hand pulse oximetry monitoring.

Infants born between 20-25 weeks gestation may be stillborn or die guickly. Initiate resuscitation and transport as soon as feasible.

If gestational age cannot be confirmed, initiate resuscitation and rapid transport.

If newborn is less than 20 weeks gestation, resuscitation is futile. Provide the newborn with warmth and consider patching to BHP for further direction.

Airway / Breath.

Cardiac/ Circula

I OC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

Medic Info.

Contact

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

APGAR Score Reference

Parameter	0	1	2
Heart rate (bpm)	0 (absent)	Slow (< 100)	<u>≥</u> 100
Respiratory effort	Absent	Slow, irregular	Good, crying
Muscle tone	None, limp	Some flexion	Active motion
Reflex irritability (suction of nares, tactile stimulation)	None	Some grimace	Good grimace, cough, cry
Colour	Blue or pale	Pink body with blue extremities	Completely pink

- ▶ APGAR performed at 1 minute & 5 minutes after delivery
- Maximum possible total score is 10 (5 parameters x maximum score 2 for each parameter)
- Don't wait for APGAR to make decision on resuscitation

Airway /

Neonatal Pre-ductal Oxygen Saturation Reference

TARGETED PRE-DUCTAL SpO2

After Birth

1 min	60-65%
2 min	65-70%
3 min	70-75%
4 min	75-80%
5 min	80-85%
10 min	85-95%

In all neonates, only apply the pulse oximeter to the **RIGHT HAND**.

Target the above values when:

- Resuscitation is anticipated
- > PPV is required for more than a few breaths
- Persistent central cyanosis, or if you need to confirm your perception of central cyanosis
- Any administration of supplemental oxygen

Destinat.

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Return of Spontaneous Circulation (ROSC) Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized

INDICATIONS

Patient with return of spontaneous circulation (ROSC) after the resuscitation was initiated

CONDITIONS

0.9% NaCl Fluid Bolus

AGE: ≥ 2 years LOA: N/A HR: N/A

RR: N/A SBP: Hypotension

Other: Chest auscultation is clear

CONTRAINDICATIONS

0.9% NaCl Fluid Bolus

Fluid overload

TREATMENT

Consider optimizing ventilation and oxygenation

Titrate oxygenation 94-98%

Avoid hyperventilation and target ETCO2 to 30-40 mmHg with continuous waveform capnography (if available)

Consider 0.9% NaCl fluid bolus (If available and authorized)

	Age	Age
	≥2 years to <12 years	≥12 years
	Route	Route
	IV	IV
Infusion	10 mL/kg	10 mL/kg
Infusion interval	Immediate	Immediate
Reassess every	100 mL	250 mL
Max. volume	1,000 mL	1,000 mL

Consider 12 lead ECG acquisition and interpretation

CLINICAL CONSIDERATIONS

Consider initating transport in parallel with the above treatment.

IV fluid bolus applies only to PCPs authorized for PCP Autonomous IV.

NOTE: Refer to page 118 for 12 Lead ECG Placement

Intro

Airway / Breath.

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced

Research/ Sp. Proj

Medical Refer.

Medic. Info

Contact

Airway / Breath

Cardiac/

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

Medic.

Contact

Cardiac Ischemia Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized.

INDICATIONS

Suspected cardiac ischemia.

CONDITIONS

ASA

AGE: ≥18 years LOA: Unaltered HR: N/A RR: N/A

SBP: N/A

Other: Able to chew and swallow

Nitroglycerin

AGE: ≥18 years LOA: Unaltered HR: 60-159 bpm RR: N/A

SBP: Normotension

Other: Prior history of nitroglycerin use

OR IV access obtained

CONTRAINDICATIONS

ASA

Allergy or sensitivity to NSAIDS
If asthmatic, no prior use of ASA
Current active bleeding

CVA or TBI in the previous 24 hours

Nitroglycerin

Allergy or sensitivity to nitrates
Phosphodiesterase inhibitor use within

the previous 48 hours
SBP drops by one-third or more of its

initial value after nitroglycerin is administered

12-lead ECG compatible with Right Ventricular MI

TREATMENT

Consider ASA

Route no

	10
Dose	160-162 mg
Max. single dose	162 mg
Dosing interval	N/A
Max. # of doses	1

Consider 12-lead ECG acquisition and interpretation for STEMI

Consider nitroglycerin

STEMI		
No	Yes	
SBP	SBP	
≥100 mmHg	≥100 mmHg	
Route	Route	
SL	SL	
0.3 OR 0.4 mg	0.3 OR 0.4 mg	
0.4 mg	0.4 mg	
5 min	5 min	
6	3	

CLINICAL CONSIDERATIONS

Dose

Max. single dose

Dosing interval Max. # of doses

Suspect a Right Ventricular MI in all inferior STEMIs and perform at minimum V4R to confirm (ST-elevation ≥ 1mm in V4R).

Do not administer nitroglycerin to a patient with Right Ventricular STEMI.

IV condition applies only to PCPs authorized for PCP Autonomous IV.

Apply defibrillation pads when a STEMI is identified.

The goal for time to 12-lead ECG from first medical contact is < 10 minutes where possible.

NOTE: Refer to page 118 for 12 Lead ECG Placement

Cardiac/Circulation Cardiac Ischemia Medical Directive

Airway / Breath.

Cardiac/ Circula

IOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer

Medic Info.

Contact

Airway / Breath

Cardiac/

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Acute Cardiogenic Pulmonary Edema Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized

INDICATIONS

Moderate to severe respiratory distress;

AND

Suspected acute cardiogenic pulmonary edema.

CONDITIONS

Nitroglycerin

AGE: ≥18 years LOA: N/A HR: 60-159 bpm

SBP: Normotension

Other: N/A

CONTRAINDICATIONS

Nitroglycerin

Allergy or sensitivity to nitrates

Phosphodiesterase inhibitor use within the previous 48 hours

SBP drops by one-third or more of its initial value after

nitroglycerin is administered



Patient Drug Dose Route Time.

Consider nitroglycerin

	SBP ≥100 mmHg to <140 mmHg IV or Hx* Yes Route SL	SBP ≥140 mmHg	
		IV or Hx*	IV or Hx*
		No Route SL	Yes Route SL
Dose	0.3 mg or 0.4 mg	0.3 mg or 0.4 mg	0.6 mg or 0.8 mg
Max. single dose	0.4 mg	0.4 mg	0.8 mg
Dosing interval	5 min	5 min	5 min
Max. # of doses	6	6	6

^{*}Hx refers to a patient with a prior history of nitroglycerin use

Consider 12-lead ECG acquisition and interpretation

CLINICAL CONSIDERATIONS

IV condition applies only to PCPs authorized for PCP Autonomous IV.

NOTE: Refer to page 118 for 12 Lead ECG Placement

Intro

Airway / Breath.

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced

Research / Sp. Proj

Medical Refer.

Medic. Info.

Contact

Airway / Breath.

Cardiac/

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Cardiogenic Shock Medical Directive - AUXILIARY

A Primary Care Paramedic may provide the treatment prescribed in this auxiliary Medical Directive if authorized for PCP Autonomous IV.

INDICATIONS

STEMI-positive 12-lead ECG;

AND

Cardiogenic shock.

CONDITIONS

0.9% NaCl Fluid Bolus

AGE: ≥18 years
LOA: N/A
HR: N/A
RR: N/A
SBP: Hypotension

Other: Chest auscultation

is clear

CONTRAINDICATIONS

0.9% NaCl fluid bolus

Fluid overload

SBP ≥90 mmHg

TREATMENT

Consider 0.9% NaCl fluid bolus Age ≥18 years Route IV Infusion 10 mL/kg Infusion interval N/A Reassess every 250 mL

1,000 mL

CLINICAL CONSIDERATIONS

Max. volume

N/A

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer.

Medic. Info.

Contact

Airway / Breath.

Cardiac/

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic.

Contact

Intravenous and Fluid Therapy Medical Directive - AUXILIARY

A Primary Care Paramedic may provide the treatment prescribed in this auxiliary Medical Directive if authorized for PCP Autonomous IV

INDICATIONS

Actual or potential need for intravenous medication **OR** fluid therapy.

CONDITIONS

IV Cannulation

AGE: ≥ 2 years
LOA: N/A
HR: N/A
RR: N/A
SBP: N/A

Other: N/A

CONTRAINDICATIONS

IV Cannulation

Suspected fracture proximal to the access site.

0.9% NaCl Fluid Bolus

AGE: ≥ 2 years LOA: N/A HR: N/A RR: N/A SBP: Hypotension

Other: N/A

0.9% NaCl Fluid Bolus

Fluid overload

TREATMENT

Consider IV cannulation

Destinat.

	Age	Age
	≥2 years to <12 years Route	≥12 years
		Route
	IV	IV
Infusion	15 mL/hr	30-60 mL/hr
Infusion interval	N/A	N/A
Reassess every	N/A	N/A
Max. volume	N/A	N/A

⚠ Mandatory Provincial Patch Point △

Patch to BHP for authorization to administer 0.9% NaCl fluid bolus to hypotensive patients ≥ 2 years to <12 years with suspected Diabetic Ketoacidosis (DKA)

Consider 0.9% NaCl fluid bolus

	Age	Age
	≥2 years to <12 years	≥12 years
	Route	Route
	IV	IV
Infusion	20 mL/kg	20 mL/kg
Infusion interval	N/A	N/A
Reassess every	100 mL	250 mL
Max. volume*	2,000 mL	2,000 mL

^{*}The maximum volume of NaCl is lower for patients in cardiogenic shock and return of spontaneous circulation.

Intro

Airway / Breath

Cardiac/

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

Medic.

Contact

Airway / Breath

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced

Research / Sp. Proi

Medical Refer.

Medic. Info

Contact

Destinat. Guide.

CLINICAL CONSIDERATIONS

"PCP Assist IV" authorizes a PCP to cannulate a peripheral IV at the request and under the direct supervision of an ACP. The patient must require a peripheral IV in accordance with the indications listed in this Medical Directive. PCPs authorized for PCP Assist IV are not authorized to administer IV fluid or medication therapy.

Microdrips and/or volume control administration sets should be considered when IV access is indicated for patients <12 years of age.

An intravenous fluid bolus may be considered for a patient who does not meet trauma TOR criteria, where it does not delay transport and should not be prioritized over management of the other reversible causes.



NOTE: Refer to page 46 for Defibrillation Joule Setting Reference Chart.



NOTE: Refer to page 120 for CPR Guidelines

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Intro

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Destinat. Guide.

Level of Consciousness/Pain/Nausea

PRIMARY CARE PARAMEDIC MEDICAL DIRECTIVES

Airway / Breath.

Hypoglycemia Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized

Cardiac/ Circula.

INDICATIONS

Suspected hypoglycemia

LOC/ Pain/ Nausea

Proced.

Research/

Sp. Proj

CONDITIONS

Dextrose

AGE: ≥ 2 years LOA: Altered

HR: N/A RR: N/A SBP: N/A

Other: Hypoglycemia

Glucagon

AGE: N/A

(≥ 4 years for IN powder)

LOA: Altered HR: N/A RR: N/A SBP: N/A

Other: Hypoglycemia

Medical Refer

> Medic. Info.

CONTRAINDICATIONS

Dextrose

Allergy or sensitivity to dextrose

Glucagon

Allergy or sensitivity to glucagon Pheochromocytoma

Contact

TREATMENT



Patient Drug Dose Route Time.

Consider glucometry

Consider **dextrose** (if available and authorized)

Age ≥ 2 years

	Concentration	Concentration
	10% dextrose	50% dextrose
	Route	Route
	IV	IV
Dose	0.2 g/kg (2 mL/kg)	0.5 g/kg (1 mL/kg)
Max. single dose	25 g (250 mL)	25 g (50 mL)
Dosing interval	10 min	10 min
Max. # of doses	2	2

Titrate dextrose to a level of awareness where the patient can safely consume complex carbohydrate.

Airway / Breath.

Cardiac/

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

Medic.

Contact

Intro

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Destinat

Consider **glucagon** (if not using dextrose) intranasal powder Age (if avthorized and N/A available) Weight Weight Weight <25 kg ≥25 kg N/A Route Route Route IM IMI IN 1 mg Dose 0.5 mg 3 mg Max. sinale 0.5 mg 1 mg 3 mg dose Dosing interval 20 min 20 min 20 min 2 2 Max. # of doses 2

CLINICAL CONSIDERATIONS

If the patient responds to dextrose or glucagon, he/she may receive oral glucose or other simple carbohydrates.

If only mild signs or symptoms are exhibited, the patient may receive oral glucose or other simple carbohydrates instead of dextrose or glucagon.

If a patient initiates an informed refusal of transport, a final set of vital signs including blood glucometry must be attempted and documented.

IV administration of dextrose applies only to PCPs authorized for PCP Autonomous IV.

Intranasal glucagon is a powder that is supplied in a commercially available single-dose intranasal device

CONSIDERATIONS FOR TREAT AND DISCHARGE (IF AUTHORIZED)

All of the following criteria must be met:

- the patient is ≥18 AND <65 years old:
- · the patient has a diagnosis of diabetes;

 the hypoglycemia can be explained by insulin administration with inadequate oral intake:

- the hypoglycemia promptly responded to a single administration of dextrose or glucagon as per the Medical Directive and/or consumed oral glucose or other complex carbohydrates;
- this was a single isolated episode of symptomatic hypoglycemia within the past 24 hours:
- the blood glucose is ≥4.0mmol/L after treatment;
- the patient has a return to their normal level of consciousness and is asymptomatic;
- · a complete set of vital signs are within expected normal ranges;

AND

- not an intentional overdose:
- the hypoglycemia must not be related to alcohol or substance abuse or withdrawal:
- · no seizure or reported history of seizure prior to paramedic treatment,
- not on an oral hypoglycemic medication:
- hypoglycemia is not considered to be related to an acute medical illness, and;
- the patient is not pregnant.

In addition to the above criteria, if all of the following requirements have been met, the patient can be discharged by paramedics:

- the patient has access to appropriate carbohydrates:
- a responsible adult agrees to remain with the patient for the next 4 hours;
- all of the patient or substitute decision makers questions were answered and a care plan was developed:
- the patient or substitute decision maker has been advised to follow up with their primary health care team or provider;
- clear instructions to call 911 were provided should symptoms redevelop;
- patient or substitute decision maker has the ability to access 911 should symptoms redevelop, and;
- · patient or substitute decision maker consents to the discharge.

CLINICAL CONSIDERATIONS (TREAT AND DISCHARGE)

Patch to BHP for consultation if you are unclear if the patient meets all of the discharge criteria.

Intro

Airway / Breath

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic.

Contact

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp.Proj

Medical Refer.

Medic. Info.

Contact

Dextrose Dosing Guide

Age	Weight kg	Blood Sugar	Dextrose prep		itial dose epeat dose	
		mmol/L		Dose g/kg	Volume ml/kg	Amt ml
< 30 days	2	< 3.0	D10W Waste 40 mls	0.2	2	4
	3		replace w/ Normal Saline		2	6
	4		Gairie		2	8
	5				2	10
≥30 days	3	< 3.0	D25W Waste 25 mls	0.5	2	6
to < 2 years	4		replace w/ Normal Saline		2	8
	5		Gairle		2	10
	6				2	12
	8				2	16
	10				2	20
	12				2	24
	14				2	28
≥ 2 years	10	< 4.0	D50W	0.5	1	10
	15				1	15
	20				1	20
	25				1	25
	30				1	30
	35				1	35
	40 45				1	40 45
	> 50				1	50

Destinat.

Nausea / Vomiting Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized.

Airway / Breath

Intro

INDICATIONS

Nausea or vomiting.

Cardiac/ Circula.

CONDITIONS

Ondansetron

AGE: N/A WEIGHT: ≥ 25 kg LOA: Unaltered HR· N/A RR: N/A SBP: N/A Other: N/A

DimenHYDRINATE

< 65 years

WEIGHT: ≥ 25 kg LOA: Unaltered HR· N/A RR: N/A SBP: N/A Other: N/A

AGE:

Proced

Research / Sp. Proj

CONTRAINDICATIONS

Ondansetron

Allergy to ondansetron

Prolonged QT symdrome (known to patient)

Apomorphine use

DimenHYDRINATE

Alleray or sensitivity to dimenHYDRINATE or other antihistamines

Overdose on antihistamines or anticholinergics or tricyclic antidepressants

Co-adminstration of diphenhydraMINE

Medical Refer.

Medic. Info

Contact

TREATMENT

Airway / Breath.



Patient Drug Dose Route Time.

Cardiac/

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

Medic.

Contact

Consider ondansetron				
	Weight			
	≥ 25 kg			
Route				
	PO			
Dose	4 mg			
Max. single dose	4 mg			
Dosing interval	N/A			
Max. # of doses	1			

Consider dimenHYDRINATE

	Weight	Weight
≥	25 kg to <50 kg	≥ 50 kg
	Route	Route
	IV/IM	IV/IM
Dose	25 mg	50 mg
Max. single dose	25 mg	50 mg
Dosing interval	N/A	N/A
Max. # of doses	1	1

CLINICAL CONSIDERATIONS

IV administration of dimenHYDRINATE applies only to PCPs authorized for PCP Autonomous IV

Prior to IV administration, dilute dimenHYDRINATE (concentration of 50 mg/1 ml) 1:9 with Normal Saline or D5W. If administered IM do not dilute

If a patient has received Ondansetron and has no relief of their nausea & vomiting symptoms after 30 minutes, dimenHYDRINATE may be considered (or vise versa.

dimenhyDRINATE can be used in patients ≥ 65 if ondansetron is not available.

Analgesia Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized

INDICATIONS

Pain

CONDITIONS

Acetaminophen

AGE: ≥12 years LOA: Unaltered HR: N/A RR· N/A

SBP: N/A Other: N/A

Ketorolac

AGE: ≥12 years LOA: Unaltered HR: N/A RR: N/A

SBP: Normotension

Other: N/A

Ibuprofen

AGE: ≥12 years LOA: Unaltered HR: N/A

RR: N/A SBP: N/A Other: N/A

Intro

Airway /

Breath

Cardiac/ Circula.

Proced

Research / Sp. Proj

Medical Refer.

Medic. Info

Contact

Airway / Breath

Cardiac/ Circula.

Proced.

Research/ Sp. Proj

Medical Refer

Medic Info.

Contact

CONTRAINDICATIONS

Acetaminophen

Acetaminophen use within previous 4 hours

Allergy or sensitivity to acetaminophen

Hy of liver disease

Active vomiting

Unable to tolerate oral medication

Suspected ischemic chest pain

Ibuprofen

NSAID use within previous 6 hours Allergy or sensitivity to ASA or NSAIDs

Patient on anticoagulation therapy

Current active bleeding

Hx of peptic ulcer disease or GI bleed Pregnant

If asthmatic, no prior use of ASA or other NSAIDs

CVA or TBI in the previous 24 hours Known renal impairment

Active vomiting

Unable to tolerate oral medication

Suspected ischemic chest pain

Ketorolac

NSAID use within previous 6 hours

Allergy or sensitivity to ASA or NSAIDs

Patient on anticoagulation therapy

Current active bleeding

Hx of peptic ulcer disease or GI bleed

Pregnant

If asthmatic, no prior use of ASA or other NSAIDs

CVA OR TBI in the previous 24 hours

Known renal impairment

Suspected ischemic chest pain

TREATMENT



Patient Drug Dose Route Time.

Airway / Breath.

Cardiac/ Circula.

Proced

Research / Sp. Proj

Medical Refer.

Medic. Info.

Contact

Destinat

Guide.

Consider acetaminophen

	Age	Age
	≥ 12 years to <18 years	≥ 18 years
Route	PO	PO
Dose	500-650 mg	960-1,000 mg
Max. single dose	650 mg	1,000 mg
Dosing interval	N/A	N/A
Max. # doses	1	1

Consider ibuprofen

	Age
	≥ 12 years
Route	PO
Dose	400 mg
Max. single dose	400 mg
Dosing interval	N/A
Max. # doses	1

Consider ketorolac

	Age
	≥ 12 years
Route	IM/IV
Dose	10-15 mg
Max. single dose	15 mg
Dosing interval	N/A
Max. # doses	1

CLINICAL CONSIDERATIONS

Whenever possible, consider co-administration of acetaminophen and ibuprofen.

Suspected renal colic patients should routinely be considered for NSAIDs, either ibuprofen or ketorolac.

IV administration of ketorolac applies only to PCPs authorized for PCP Autonomous IV.

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

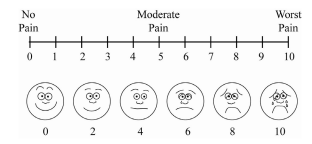
Medic. Info.

Contact

Destinat. Guide.

Pain Scale Reference

Can be utilized for patients 3 years of age and older



Opioid Toxicity and Withdrawal Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized.

INDICATIONS

Suspected opioid toxicity.

CONDITIONS

Naloxone

AGE: ≥ 24 hours LOA: Altered HR: N/A

RR: <10 breaths/min

SBP: N/A

Other: Inability to adequately ventilate **OR** persistent need to assist ventilations

burprenorphine/naloxone

AGE: ≥ 16 LOA: Unaltered

HR: N/A RR: N/A SBP: N/A

Other: Received naloxone for current

opioid toxicity episode

Patient is exhibiting acute withdrawal with a COWS* score > 8

CONTRAINDICATIONS

Naloxone

Allergy or sensitivity to naloxone

buprenorphine/naloxone

Allergy or sensitivity to buprenorphine

Taken methadone in the past 72

hours

Intro

Airway / Breath

Cardiac/ Circula.

> OC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic.

Contact

Destinat.

TREATMENT

Airway / Breath.



Patient Drug Dose Route Time.

Cardiac/ Circula.

Proced.

Research/ Sp. Proj

Medical Refer

Medic Info.

Contact

Consider naloxone

	Route	Route	Route	Route
	IV	IM	IN	SC
Dose	Up to 0.4 mg*	0.4 mg	2-4 mg	0.8 mg
Max. single dose	0.4 mg	0.4 mg	2-4 mg	0.8 mg
Dosing interval	5 min	5 min	5 min	5 min
Max. # of doses	3	3	3	3

*For the IV route, titrate naloxone only to restore the patient's respiratory status.

Consider buprenorphine/naloxone (if available and authorized)

Route BUC/SL Initial dose 16 mg Subsequent 8 mg dose(s) dose Dosing interval 10 minutes Max. cumulative 24 mg dose

CLINICAL CONSIDERATIONS

IV administration of naloxone applies only to PCPs authorized for PCP Autonomous IV.

Upfront aggressive management of the airway is paramount and the initial priority.

If no response to initial treatment; consider patching for further doses.

If the patient does not respond to airway management and the administration of naloxone, glucometry should be considered.

Combative behaviour should be anticipated following naloxone administration and paramedics should protect themselves accordingly, thus the importance of gradual titrating (if given IV) to desired clinical effect: respiratory rate ≥10, adequate airway and ventilation, not full alertness.

Intro

Airway / Breath

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer.

Medic.

Contact

Airway / Breath.

*Clinical Opiate Withdrawal Scale (COWS)

< 5 - No active withdrawal		< 36 - Severe withdrawal
5-12 - Mild withdrawal	25-36 - Moderately severe withdrawal	Williurawai

A score of ≥8 is an indication for buprenorphine/naloxone administration

Cardiac/ Circula.

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

GI Upset over last ½ hour 0 no of isymptoms 1 stomach cramps 2 nausea or loose stool 3 vomiting or diarrhea 5 multiple episodes of diarrhea or vomiting
Tremor observation of outstretched hands 0 no tremor 1 tremor can be felt, but not observed 2 slight tremor observable 4 gross tremor or muscle twitching
Yawning observation during assessment 0 no yawning 1 yawning once or twice during assessment 2 yawning three or more times during assessment 4 yawning several times/minute
Anxiety or Irritability 0 none 1 patient reports increasing irritability or anxiousness 2 patient obviously irritable anxious 4 patient so irritable or anxious that participation in the assessment is difficult
Gooseflesh Skin 0 skin is smooth 3 pilloerrection of skin can be felt or hairs standing up on arms 5 prominent piloerrection
Total Score The total score is the sum of all 11 items

Suspected Adrenal Crisis Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized.

INDICATIONS

A patient with primary adrenal failure who is experiencing clinical signs of an adrenal crisis.

CONDITIONS

Hydrocortisone

AGE: N/A
LOA: N/A
HR: N/A
RR: N/A
SBP: N/A

Other: Paramedics are presented with a vial of hydrocortisone for the identified

patient AND

Age-related hypoglycemia OR

GI symptoms (vomiting, diarrhea, abdominal pain) OR

Syncope OR

Temperature ≥38C or

suspected/history of fever **OR** Altered level of awareness **OR**

Age-related tachycardia OR

Age-related hypotension

Intro

Airway / Breath

Cardiac/ Circula.

> OC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer.

Medic.

Contact

CONTRAINDICATIONS

hydrocortisone

Airway / Breath.

Circula.

Hydrocortisone
Allergy or sensitivity to

Cardiac/

TREATMENT



Patient Drug Dose Route Time.

Consider hydrocortisone

Route
IM/IV

Dose 2 mg/kg*

Max. single dose 100 mg

Dosing interval N/A

Max. # doses 1

*Dose should be rounded to the nearest 10 mg

Medical Refer

Research/ Sp. Proj

Proced.

CLINICAL CONSIDERATIONS

 $\ensuremath{\mathsf{IV}}$ Administration of hydrocortisone applies only to PCP's authorized for PCP Autonomous $\ensuremath{\mathsf{IV}}.$

Medic. Info.

Contact

Seizure Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Auxiliary Medical Directive if authorized.

CONSIDERATIONS FOR TREAT AND DISCHARGE (IF AUTHORIZED)

All of the following criteria must be met:

- the patient is ≥18 AND <65 years old;
- · patient must have a history of epilepsy;
- the patient is taking their anticonvulsant medication as prescribed;
- the patient must have only had a single seizure episode in the past 24 hours;
- the seizure pattern and duration must be similar to past seizures:
- the patient has returned to their normal level of consciousness:
- a complete set of vital signs including temperature are within expected normal ranges;

AND

- the seizure must not be related to hypoglycemia, alcohol or substance abuse or withdrawal:
- the patient must not have received midazolam by paramedics;
- the patient did not injure themselves during seizure activity;
- the patient must not have a fever, preceding illness or recently started a new medication, and:
- the patient is not pregnant.

In addition to the above criteria, if all of the following requirements have been met, the patient can be discharged by paramedics:

- a responsible adult agrees to remain with the patient for the next 4 hours;
- all of the patient or substitute decision makers questions were answered and a care plan was developed:
- the patient or substitute decision maker has been advised to follow up with their primary health care team or provider;
- · clear instructions to call 911 were provided should symptoms redevelop;
- patient or substitute decision maker has the ability to access 911 should symptoms redevelop, and
- patient or substitute decision maker consents to the discharge.

CLINICAL CONSIDERATIONS (TREAT AND DISCHARGE)

Patch to BHP for consultation if you are unclear if the patient meets all of the discharge criteria.

Intro

Airway / Breath

Cardiac/ Circula.

> .OC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer.

Medic.

Contact

Destinat.

Intro

Airway /
Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Destinat. Guide.

Intentionally Left Blank

Procedural

PRIMARY CARE PARAMEDIC MEDICAL DIRECTIVES



Intro

Airway / Breath.

Home Dialysis Emergency Disconnect Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized.

Cardiac/ Circula.

LOC/ Pain/

Nausea

INDICATIONS

Patient receiving home dialysis (hemo or peritoneal) and connected to dialysis machine and requires transport to the closest appropriate receiving facility;

AND

Patient is unable to disconnect:

AND

There is no family member or caregiver who is available and knowledgeable in dialysis disconnect.

Proced

Research /

Sp. Proj

Medical

Refer.

CONDITIONS

Home Dialysis Emergency Disconnect

AGE: N/A LOA: N/A HR: N/A RR: N/A

Other: N/A

Medic. Info.

CONTRAINDICATIONS

Home Dialysis Emergency Disconnect

N/A

Contact

TREATMENT

Consider Home Dialysis Emergency Disconnect

CLINICAL CONSIDERATIONS

Generally, emergency disconnect kit with materials and instructions can be found hanging from dialysis machine or nearby on the wall.

Ensure both the patient side and machine side of the connection are clamped <u>before</u> disconnecting and attaching end caps.

Intro

Airway / Breath

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced

Research/ Sp. Proj

Medical Refer.

Medic.

Contact

Destinat.

Emergency Dialysis Disconnect Prompt Card

Airway / Breath

Cardiac/

Circula.

Hemodialysis Disconnect

- Clamp patient side tubing clamps
- Clamp machine side clamps
- Attach sterile Luer Lock caps to the ends of the patient tubing
- Disregard any alarms that may sound from the machine
- Secure patient tubing and cover with abdo pad

IOC/ Pain/ Nausea

Continuous Ambulatory Peritoneal Dialysis (CAPD)

- Close the twist clamp
- Clamp both the fill and drain bag tubing with clamps supplied in disconnect kits
- Screw a sterile Luer Lock on the patient side tubing
 - · Snap a sterile Luer Lock on the machine side tubing
- Secure patient tubing and cover with abdo pad

Research / Sp. Proj

Medical Refer

Medic

Info.

Automatic Peritoneal Dialysis (APD)

- Push "Stop" button on ADP machine
- Close the twist clamp
- Disconnect the patient tubing from the machine tubing
- Screw a sterile mini cap on the patient tubing
- Snap a mini cap on the machine tubing
- Secure patient tubing and cover with abdo pad

Contact

Emergency Childbirth Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized.

INDICATIONS

Pregnant patient experiencing labour; OR

Post-partum patient immediately following delivery and/or placenta.

CONDITIONS

Delivery

AGE: Childbearing years

LOA: N/A HR: N/A RR: N/A SBP: N/A

Other: Second stage labour AND/OR

Imminent birth AND/OR Shoulder Dystocia AND/OR Breech Delivery AND/OR Prolapsed Cord

External Uterine Massage

AGE: Childbearing years

LOA: N/A HR: N/A RR: N/A SBP: N/A

Other: Post-placental delivery

Umbilical Cord Management

AGE: Childbearing years

LOA: N/A HR: N/A RR: N/A SBP: N/A

Other: Cord complications

OR

if neonatal or maternal resuscitation is required

OR

Due to transport considerations

Oxytocin

AGE: Childbearing years

LOA: N/A HR: N/A RR: N/A

SBP: < 160 mmHg Other: Postpartum delivery

AND/OR
Placental delivery

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer

Medic.

Contact

Destinat.

Airway / Breath

Cardiac/

Circula.

IOC/

Pain/ Nausea

CONTRAINDICATIONS

Delivery

N/A

Umbilical Cord Management

External Uterine Massage

Placenta not delivered

Oxytocin

Allergy or sensitivity to oxytocin

Undelivered fetus

Suspected or known pre-eclampsia with current pregnancy

Eclampsia (seizures) with current pregnancy

≥4 hours post placenta delivery

Research/ Sp. Proj

Medical Refer

Medic

Info.

TREATMENT

Consider delivery

Position the patient and deliver neonate.

Consider shoulder dystocia delivery

Perform ALARM twice on scene. If successful; deliver neonate. If unsuccessful; transport to closest appropriate facility

Consider breech delivery

HANDS OFF the breech. Allow neonate to deliver to umbilicus; consider carefully releasing the legs & arms as they are delivered; otherwise hands off.

Once hairline is visible **AND/OR** 3 mins has passed since umbilicus was visualized attempt the Mauriceau Smellie-Veit maneuver.

If successful; deliver neonate. If unsuccessful; transport to closest appropriate facility.

Contact

Consider prolapsed cord delivery

If a cord prolapse is present, the fetal part should be elevated to relieve pressure on the cord. Assist the patient into a knee-chest position or exaggerated Sims position, and insert gloved fingers/hand into the vagina to apply manual digital pressure to the presenting part which is maintained until transfer of care in hospital.

Consider umbilical cord management

If a nuchal cord is present and loose, slip cord over the neonate's head. Only if a nuchal cord is tight and cannot be slipped over the neonate's head, clamp and cut the cord, encourage rapid delivery.

Following delivery of the neonate, the cord should be clamped and cut immediately if neonatal or maternal resuscitation is required. Otherwise, after pulsations have ceased (approximately 2-3 minutes), clamp the cord in two places and cut the cord.

Consider external uterine massage

Post placental delivery

Consider oxytocin (where authorized and available)

	Route	
	IM	
Dose	10 units	
Max. single dose	10 units	
Dosing interval	N/A	
Max. # of doses	1	

Airway / Breath

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced

Research/ Sp. Proj

Medical Refer

Medic.

Contact

CLINICAL CONSIDERATIONS

Airway / Breath

If the patient presents with limb-presentation, do not attempt to push the limb back into the vagina; discourage the patient from pushing, cover the limb using a dry sheet to maintain warmth, and initiate transport as per the Load and Go Patient Standard of the BLS PCS

Cardiac/ Circula.

If labour is failing to progress, discourage the patient from pushing or bearing down during contractions.

IOC/ Pain/ Nausea If delivery has not occurred at scene within approximately ten minutes of initial

- a. Patient assessment findings:
 - i. Lack of progression of labour:

assessment, consider transport in conjunction with the following:

- ii. Multiple births expected:
- iii. Neonate presents face-up:
- iv. Pre-eclampsia;
- v. Presence of vaginal hemorrhage;
- vi. Premature labour;
- vii. Primip:
- Distance to the closest appropriate receiving facility.

When the placenta is delivered, inspect it for wholeness, place in a plastic bag from the OBS kit, label it with the maternal patient's name and time of delivery. and transport it with the maternal or neonatal patient. Delivery of the placenta should not delay transport considerations/initiation.

Research / Sp. Proj

Medical Refer

Medic Info.

Contact

Intro Airway / Breath. Cardiac/ Circula. Nausea

LOC/ Pain/

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Contact

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Research / Special Projects PRIMARY CARE PARAMEDIC MEDICAL DIRECTIVES



Intro

Airway / Breath

Special Project Palliative Care Medical Directive

A Primary Care Paramedic may provide the treatment and/or patient disposition prescribed in this Medical Directive if authorized.

Patch

If a paramedic determines that the patient would benefit from any other management that is not included in this special project medical directive, a patch to a BHP is necessary.

Registered Patient

A registered patient is under the care of a palliative care team through Home and Community Care, or a physician or nurse practitioner providing palliative care services in the community. The paramedic is required to confirm the patient registration according to their local process.

Management of Patients with Palliative Care Needs

Patients with palliative care needs may require a different approach to assessment and treatment that reflects their unique goals of care. Therefore paramedics, for this defined patient population, should consider prioritizing patient comfort and are not required to follow the described regimen of strict vital signs, cardiac monitoring and transport as directed in the Basic Life Support Patient Care Standard (BLS PCS). If patient transport is initiated, however, paramedics should consider usual care (vitals and monitoring) per the ALS and BLS PCS in conjunction with the patient's goals of care; they may also consider symptom treatments below if indicated.

Medical Directive

This Medical Directive is written in five sections or equivalent to five directives combined including four symptom-based sections (Dyspnea, Hallucinations/Agitation, Nausea/Vomiting and Terminal Congested Breathing) as well as a Treat and Refer directive. Any of these directives can apply, individually or in combination, to a patient with palliative care needs. The Treat and Refer part of this directive can be applied even if no symptoms listed in the directive are present or treatments have not been provided. All patients who

Cardiac/

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

remain at home must be referred to their palliative care team to ensure follow up of their presenting complaint.

When in doubt, please consult/patch to a Base Hospital Physician (BHP) in consultation with palliative physician or nurse if available.

Breath.

Airway /

Cardiac/

Circula.

LOC/ Pain/

Nausea

DYSPNEA

INDICATIONS

Registered Palliative Care Patient

And

Uncontrolled dyspnea with suspected bronchoconstriction

CLINICAL CONSIDERATIONS

Salbutamol should only be used in patients whose dyspnea is accompanied by wheezing or a history of response to bronchodilators. Proced.

Salbutamol

AGE: ≥18 LOA: N/A HR: N/A RR: N/A SBP: N/A

CONDITIONS

Other: For Dyspnea with suspected bronchoconstriction

only

Medical Refer

Medic.

Contact

Intro

Airway / Breath. CONTRAINDICATIONS

Salbutamol

Allergy to salbutamol

Cardiac/ Circula.

IOC/

Pain/ Nausea

Proced.

TREATMENT

5Rs

Patient Drug Dose Route Time.

Consider Salbutamol

Route Route NEB MDI* Up to 800 mcg 5 mg Dose (8 puffs) 5mg Max. dose 800 mcg 5-15 min prn Dosing interval 5-15 min prn 3 Max. # of doses

*1 puff - 100 mcg

Medical Refer

Medic

Info.

HALLUCINATIONS OR AGITATION

INDICATIONS

Registered Palliative Care Patient

And

Increasing agitation or suspected new or increased hallucinations

Contact

CLINICAL CONSIDERATIONS

Destinat.

Research / Sp. Proj. Special Project Palliative Care Medical Directive

103

CONDITIONS

Haloperidol

AGE: ≥18

LOA: N/A

HR: N/A

RR: N/A

SBP: N/A

Other: N/A

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

. . .

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Medical Refer

Medic.

Contact

CONTRAINDICATIONS

Haloperidol

Allergy to haloperidol

Known Parkinson's or Lewy Body Dementia

Neuroleptic Malignant Syndrome

Destinat.

TREATMENT

Airway / Breath.



Patient Drug Dose Route Time.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Consider Haloperidol

	Route
	SC
Dose	0.5-1 mg
Max. single dose	1 mg
Dosing interval	30 min
Max. # of doses	2

Research/

NAUSEA OR VOMITING

INDICATIONS

Medical Refer Registered Palliative Care Patient

And

Nausea and/or vomiting

Medic. Info.

CLINICAL CONSIDERATIONS

Contact

Dimenhydrinate is rarely used in the palliative care population as it can cause delirium, increase drowsiness, and does not target the appropriate receptors to control the nausea in most patients. It should only be used in patients with contraindications to haloperidol where ondansetron cannot be used and should be started at low doses.

Destinat. Guide.

Research / Sp. Proj. Special Project Palliative Care Medical Directive

CONDITIONS

Haloperidol	Ondansetron	Dimenhydrinate
AGE: ≥18	AGE: ≥18	AGE: ≥18
LOA: N/A	LOA: N/A	LOA: N/A
HR: N/A	HR: N/A	HR: N/A
RR: N/A	RR: N/A	RR: N/A
SBP: N/A	SBP: N/A	SBP: N/A
Other: N/A	Other: Contraindication to Haloperidol	Other: Contraindication to Haloperidol

CONTRAINDICATIONS

Haloperidol	Ondansetron
Allergy to haloperidol	Allergy to ondansetron
Known Parkinson's or Lewy Body Dementia	
Neuroleptic Malignant Syndrome	

Dimenhydrinate

Allergy to dimenhydrinate or other antihistamines

Overdose on antihistamines or anticholinergics or tricyclic antidepressants

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/

Medical Refer.

Medic. Info.

Contact

Destinat.

TREATMENT

Airway / Breath.



Patient Drug Dose Route Time.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research Sp. Proj

Medical Refer.

Medic. Info.

Contact

Destinat.

Consider Haloperidol

	Route
	SC
Dose	0.5-1 mg
Max. single dose	1 mg
Dosing interval	30 min
Max. # of doses	2

Consider Ondansetron

	Route
	PO/SC
Dose	4 mg
Max. single dose	4 mg
Dosing interval	N/A
Max. # of doses	1

Research / Sp. Proj. Special Project Palliative Care Medical Directive

Consider Dimenhydrinate

	Route
	SC
Dose	25-50 mg
Max. single dose	50 mg
Dosing interval	N/A
Max. # of doses	1

TERMINAL CONGESTED BREATHING

INDICATIONS

Registered Palliative Care Patient

And

Congested/loud/rattling breathing in patients near the end of life

CLINICAL CONSIDERATIONS

Patient repositioning and gentle turning of the head to the side can be done instead of medication however suction of the oropharynx is not appropriate as it will likely cause discomfort and a gag reflex. Airway /

Breath

Cardiac/

LOC/ Pain/ Nausea

Proced

Research/ Sp. Proi

Medical Refer.

Medic.

Contact

CONDITIONS

Airway / Breath.

Cardiac/ Circula. Glycopyrrolate or Atropine

AGE: ≥18 LOA: N/A

HR: N/A RR: N/A

SBP: N/A Other: N/A

LOC/ Pain/ Nausea

CONTRAINDICATIONS

GlycopyrrolateAllergy to glycopyrrolate

Atropine Allergy to atropine

Proced.

Research/

Medical Refer

TREATMENT

Medic.

Rs

Patient Drug Dose Route Time.

Contact

Destinat. Guide.

 ${\sf Research\,/\,Sp.\,Proj.}\ \ \textbf{Special\,Project\,Palliative\,Care\,Medical\,Directive}$

Consider Glycopyrrolate or Atropine

	Route
	SC
Dose	0.4 mg
Max. single dose	0.4 mg
Dosing interval	N/A
Max. # of doses	
	4
	1

TREAT AND REFER

INDICATIONS

Registered Palliative Care Patient

And

Symptoms improved to patient's/Substitute Decision Maker's (SDM) satisfaction

And

After informed discussion patient/SDM preference to remain at home

CLINICAL CONSIDERATIONS

- A period of observation is recommended after the administration of any medication if the patient is not transported to ensure adequate response and no unexpected immediate adverse effects
- Transport should be considered if there is strong suspicion of reversible causes including but not limited to:
 - o Complete bowel obstruction with no prior history of same
 - New Spinal Cord Compression
 - New Superior Vena Cava (SVC) Obstruction
 - Airway obstruction
 - Suspected new pathologic fracture
- If patients do not meet the treat and refer conditions, paramedics should consider consulting BHP, follow the patient refusal standard and document appropriately.

Intro

Airway / Breath

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced

Research/ Sp. Proi

Medical Refer.

Medic.

Contact

Destinat.

Intro

CONDITIONS

Airway / Breath Age ≥ 18

DNR and/or previous goals of care discussion

Registered Palliative Care Patient

Cardiac/ Circula.

IOC/

Pain/ Nausea CONTRAINDICATIONS

Concerns of patient abuse or neglect

Patient and SDM cannot demonstrate decision-making capacity based on the

Aid to Capacity Evaluation Tool

Uncontrolled or new seizures

TREATMENT

Paramedics may assess and/or treat patients according to this medical directive and, in collaboration with the patient/SDM, honour wishes to remain at home (treat and refer). Paramedics will notify the patient's palliative care team for all patients who remain at home to ensure follow up for their presenting complaint.

Proced.

Research/ Sp. Proj

Medical Refer

Medic.

Contact

Research Study Medical Directive for Palliative Care (E3CP)

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if specifically authorized.

INDICATIONS

A patient, or their surrogate, who self identifies as receiving palliative care and who is experiencing the following uncontrolled symptoms:

- pain
- nausea
- delirium or agitation
- dyspnea
- noisy breathing or excessive secretions

CONDITIONS

Haloperidol	Glycopyrrolate
AGE: ≥ 18 years	AGE: ≥ 18 years
LOA: n/a	LOA: n/a
HR: n/a	HR: n/a
SBP: n/a	SBP: n/a
Delirium, agitation, or nausea	Other: Excessive secretions or noisy breathing

CONTRAINDICATIONS

Haloperidol	Glycopyrrolate
Allergy or sensitivity to haloperidol	Allergy or sensitivity to glycopyrrolate
History of Parkinson's Disease, Lewy Body	
Dementia, or extrapyramidal symptoms	
from medications	

TREATMENT

Consider Haloperidol		
	Indication	
	Nausea, delirium, or agitation	
	Route	
	SC/IV	
Dose	0.5-1 mg	
Max. single dose	1 mg	
Dosing interval	30 minutes	
Max # of doses	2	

Intro

Airway / Breath.

Cardiac/

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proi

Medical Refer.

Medic. Info.

Contact

Destinat.

Intro

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research Sp. Proi

Medical Refer

Medic. Info.

Contact

Destinat.

Consider Glycopyrrolate				
	Indication Secretions or noisy breathing Route SC			
Dose	0.4 mg			
Max. single dose	0.4 mg			
Dosing interval	n/a			
Max # of doses	1			

!! Local Mandatory Patch Point !!

- 1. Patch to the BHP if patient symptoms not controlled with medical directives.
- 2. Patch to the BHP if patient goals of care are unclear.
- 3. Patch to the BHP for all non-transport situations.

Intro

Airway / Breath

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced

Sp. Proj

Medical Refer.

Medic. Info

Contact

Destinat

Guide.

Research / Sp. Proi Study Medical Directive for Palliative Care Symptom Relief Subcutaneous Line Placement Medical Directive

114

A Primary Care or Advanced Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized. This directive is to be used only in conjunction with Study Medical Directive for Palliative Care Symptom Relief (EC3P).

Study Medical Directive for Palliative Care Symptom Relief - Subcutaneous Line Placement

Medical Directive

INSERTION OF SUBCUTANEOUS LINE

INDICATIONS

A patient, or their surrogate, who self identifies as palliative and is being treated under the Study Medical Directive for Palliative Care Symptom Relief by Paramedics

And

Parenteral administration of palliative care symptom relief medications is clinically indicated (such as Morphine, Hydromorphone, Haloperidol, Midazolam)

And

It is expected more than one medication administration will be required and thus the patient will benefit from placement of a subcutaneous line

And

A follow up plan is in place to ensure ongoing management of the subcutaneous line (such as follow up by MRP or community paramedic)

CONTRAINDICATIONS

N/A

TREATMENT



Patient Drug Dose Route Time.

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced.

Medical Refer

Medic Info.

Contact

Destinat Guide.

Consider Subcutaneous Line Placement

CLINICAL CONSIDERATIONS

- A period of observation is recommended after the administration of any medication if the patient is not transported to ensure adequate response and no unexpected immediate adverse effects
- · Adverse events after insertion include pain at the site (from the irritation of the drug or the injection was fast, this is prevented by injecting the drug slowly). If pain remain then the needle may be pulled back into the intradermal space (put a folded 2x2 gauze under the butterfly wings to elevate the needle to 45 degrees. If pain persist, then you need to change needle).

Research / Sp. Proj Study Medical Directive for Palliative Care Symptom Relief Subcutaneous Line Placement Medical Directive

The PRIME Trial Medical Directive

A Primary Care Paramedic may provide the treatment prescribed in this Medical Directive if authorized.

Indications

Pediatric non-traumatic cardiac arrest

Conditions

	CPR
Age	≥ 24 hours to 17 years
LOA	Altered
HR	N/A
RR	N/A
SBP	N/A
Other	Performed in 2 minute intervals

	Manual Defibrillation
Age	≥ 24 hours to 17 years
LOA	Altered
HR	N/A
RR	N/A
SBP	N/A
Other	VF OR pulseless VT

	AED Defibrillation
Age	≥ 24 hours to 17 years
LOA	Altered
HR	N/A
RR	N/A
SBP	N/A
Other	Defibrillation indicated

Epinephrine				
Age	≥ 24 hours to 17 years			
LOA	Altered			
HR	N/A			
RR	N/A			
SBP	N/A			
Other	N/A			

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/

Medical Refer.

Medic.

Contact

Intro

Airway / Breath.

CPR

Obviously dead as per BLS PCS

Meet conditions of Do Not Resuscitate (DNR) Standard

Contraindications

Manual Defibrillation

Rhythms other than VF or pulseless VT

Cardiac/ Circula.

IOC/ Pain/ Nausea AED Defibrillation

Non-shockable rhythm

Epinephrine

Allergy or sensitivity to epinephrine

Treatment

Proced.

Consider CPR as described in the BLS PCS

Medical Refer

Medic Info.

Contact

Consider manual defibrillation (if available and authorized) Age Age ≥ 24 hours to < 8 years ≥ 8 years to 17 years Dose 1 defibrillation 1 defibrillation Initial dose 2 J/kg As per BH / manufacturer Subsequent dose(s) 4 J/kg As per BH / manufacturer Dosing interval 2 min 2 min Max. # of doses N/A N/A

Consider AED defibrillation (if not using manual defibrillation)					
	A	Age			
	≥ 24 hours to < 8 years		≥ 8 years to 17 years		
	With Pediatric Attenuator Cable	Without Pediatric Attenuator Cable	N/A		
Dose	1 defibrillation 1 defibrillation		1 defibrillation		
Max. single dose	As per BH / manufacturer	As per BH / manufacturer	As per BH / manufacturer		

2 min

N/A

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Consider EPINEPHrine Preload (if available)

Dosing interval

Max. # of doses

Route

2 min

N/A

2 min

N/A

IM

	Weight				
	\geq 3 kg to \leq 5 kg	≥ 5 kg to < 10 kg	≥ 10 kg to < 20 kg	≥ 20 kg to < 30 kg	≥ 30 kg
Dose	0.3 mg	0.5 mg	1 mg	2 mg	3 mg
Total # of injections	1	1	1	1	1
Dosing interval	N/A	N/A	N/A	N/A	N/A
Max. # of doses	1	1	1	1	1

Proced.

Research/

Medical Refer.

Medic. Info.

Contact

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp.Proj

Medical Refer

Medic.

Contact

Destinat. Guide.

Consider EPINEPHrine

Route

IM

		Weight		
	\geq 3 kg to \leq 5 kg	\geq 5 kg to < 10 kg	$\geq 10~kg$ to $<20~kg$	≥ 20 kg
Dose	0.3 mg	0.5 mg	1.0 mg	2 mg
IM autoinjector used	0.3 mg	0.5 mg	0.5 mg	0.5 mg
Total # of injections	1	1	2	4
Dosing interval	N/A	N/A	N/A	N/A
Max. # of doses	1	1	1	1

Clinical Considerations

IM epinephrine to be administered as soon as feasible after the initial analysis is completed by paramedics.

Continue standard care as per the Medical Cardiac Arrest Directive once IM epinephrine has been administered.

Medical References

PRIMARY CARE PARAMEDIC MEDICAL DIRECTIVES

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

Medic. Info.

Contact

Destinat. Guide.

ETCO₂ Waveforms

Sudden loss waveform

- ET tube disconnected, dislodged, kinked or obstructed
- Loss of circulatory function



Decreasing EtC2

- ET tube cuff leak
- ET tube in hypopharynx
- Partial obstruction



CPR Assessment

 Attempt to maintain minimum of 10 mmHg



Sudden increase in EtCO2

Return of spontaneous circulation (ROSC)



Bronchospasm

("Shark-fin" appearance)

- Asthma
- COPD



Hypoventilation



Hyperventilation

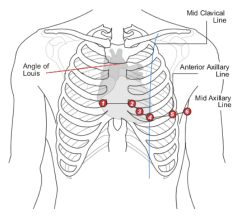


Decreased EtCO2

- Apnea
- Sedation



12 Lead ECG Placement



PRECORDIAL LEADS:

V1 - 4th intercostal space to the right of the sternum

V2 - 4th intercostal space to the left of the sternum

V3 - directly between leads V2 and V4

 ${
m V4}\,$ - $5^{
m th}$ intercostal space at left midclavicular line

V5 - level with lead V4 at left anterior axillary line

V6 - level with lead V5 at left midaxillary line

LIMB LEADS

RA - right forearm or wrist

LA - left forearm or wrist

LL - left lower leg

RL - right lower leg

NOTE:

Refer to the Medical Directives for the clinical situations where a 12-Lead ECG should be considered. This may include patients experiencing cardiac ischemia, acute cardiogenic pulmonary edema, tachycardias, bradycardias, shortness of breath or upon ROSC.

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic.

Contact

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

Medic. Info.

Contact

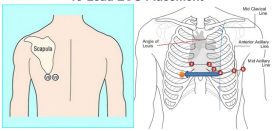
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STEMI Anatomical Location

I	aVR	V1	V4
Lateral		Septal	Anterior
Ш	aVL	V2	V5
Inferior	Lateral	Septal	Lateral
Ш	aVF	V3	V6
Inferior	Inferior	Anterior	Lateral

15-Lead ECG Placement



V4 becomes V

V6 becomes

V4R - fifth intercostal space at **right** midclavicular line (similar position as V4 but on right side of

V5 becomes V8 - level v

V9

level with V6 at left midscapular line
 level with V6 at left paravertebral line

NOTE:

- Limb leads should be placed on the limbs and not on the chest
- Consider assessing V4R when the 12 Lead identifies an inferior STEMI or ST depression in any of the septal leads (V1/V2).
- Consider assessing V8 and V9 when the 12 lead shows ST depression in the precordial leads or the 12 lead appears 'normal'.
- ST elevation of ≥ 1 mm in V4R and inferior ST-elevation, suggests a Right Ventricular involvement.
- ST elevation of \geq 1 mm or greater in V8 and V9 suggests Posterior MI.

CPR Guidelines

		Recommendations				
Component	★ Adults	* Children	★ Infants			
Recognition	★★★ Check for responsiveness (for all ages) ★★★ No breathing or only gasping (ie, abnormal) ★★★ No pulse palpated within 10 seconds for all ages ★★ HR < 60 and signs of hypoperfusion					
CPR sequence	★★★ C-A-B					
Compression rate	★★★ 100-120/r	nin				
Compression depth	★ 5.0 – 6.0 cm (2.0 - 2.4 inches)	★ At least ¹ / ₃ AP diameter ★ About 5 cm (2 inches)	★ At least ¹/₃ AP diameter ★ About 4 cm (1¹/₂ inches)			
Chest wall recoil	*** Allow complete recoil between compressions Rotate compressors every 2 minutes					
Compression interruptions		interruptions in ches o limit interruptions				
Airway	★★★ Head tilt-cl jaw thrust	hin lift or where trau	ma is suspected,			
Compression-to- ventilation ratio (until advanced airway placed)	★ 30:2 1 or 2 rescuers	Single res ★★ 15:2	Single rescuer ★★ 15:2 2 HCP rescuers			
Ventilations with advanced airway (HCP)	Neonates: 3:1 1 breath every 6-8 seconds (10 breaths/min) Asynchronous with chest compressions About 1 second per breath without too much force Visible chest rise					
Defibrillation	Visible chest rise ★★★ Attach and use AED as soon as available. Minimize interruptions in CPR pre & post rhythm interpretation/defibrillation to < 10 seconds					

CPR NOTES:

- ▶ Rate: 100-120 compressions/minute and allow full chest recoil.
- Switch person doing compressions every 2 minutes and focus on high quality CPR.
- Minimize interruntions to cheet compressions at all times

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ADULTS:

Airway / Breath Non-intubated: ratio 30:2 as above.

Intubated: 10 ventilations per minute without interrupting chest compressions. SGA inserted: 10 ventilations per minute without interrupting chest compressions

PEDIATRICS (30 DAYS TO AGE 12):

Non-intubated: ratio 15:2 as above.

Intubated: 10 ventilations per minute without interrupting chest compressions. Ventilations for resp. arrest only, non-intubated: 12-20/min.

NEONATE:

Non-intubated AND intubated 3:1 ratio as above.

ETCO2 IN CARDIAC ARREST

- When a SGA or ETT is in place, the following concepts apply:
- Continuous waveform capnography is recommended in addition to clinical assessment as the most reliable method of confirming and monitoring correct placement of an endotracheal tube
- Waveform capnography should be used to confirm and monitor endotracheal tube and SGA placement at all times
- Studies on waveform capnography have shown nearly 100% sensitivity and 100% specificity in identifying correct endotracheal tube and SGA placement
- Using quantitative waveform capnography is recommended in patients to monitor CPR quality, optimize chest compressions, and detect ROSC during chest compressions or when rhythm check reveals an organized rhythm (in addition to pulse checks)
- If waveform capnography abruptly increases to a normal value (35 to 40 mm Hg or higher) and is sustained, this may represent ROSC; wait for the next rhythm check to check for a pulse (or stop sooner if the patient exhibits signs of life)
- An ETCO₂ < 10 mmHg in VSA patients after 20 minutes of ACLS have a very poor prognosis; and can be used with clinical factors for the BHP to determine if TOR is appropriate.

LOC/

Nausea

Circula.

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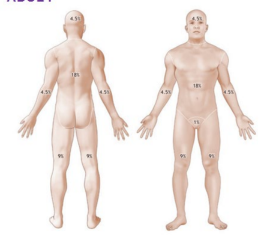
Contact

Rule of Nines, Burn Percentage Chart

PEDIATRIC



ADULT



Advanced Trauma Life Support, 9th Edition 2012 ; The American College of Surgeons.

Medical References Rule of Nines, Burn Percentage Chart

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Intramuscular Injection

- An intramuscular (IM) injection is a parenteral medication administration route commonly used by paramedics. It involves injecting a pharmacological agent directly into muscle tissue. Muscle tissue is very vascular, and as a result IM injections tend to have a faster onset of action than subcutaneous administrations.
- Identify patient that meets criteria for an intramuscular medication administration (refer to the Medical Directives or BHP order).
- ▶ Ensure all the "rights" of medication administration have been met
- ▶ Confirm medication and dose with paramedic partner if available.
- ▶ Follow safe process for responsible medication administration.
- Landmark the intended injection site. Generally the deltoid and the vastus lateralis are easily accessible and appropriate sites for IM injections; however other sites may be appropriate and can be landmarked as per the diagram on the following page.
- Select the appropriate size and gauge needle.
- Cleanse the needle insertion site using aseptic technique.
- Prepare the appropriate medication and dose into the syringe and needle ensuring all air bubbles are removed prior to injection.
- Stretch the skin taut and use the "Z-track" technique to displace the skin and soft tissue. Insert the needle with syringe/medication at a 90 degree angle using a "dart style" motion. The Z-track method reduces the chance the medication will leak from the muscle into the subcutaneous tissue.
- Inject the correct dose of medication.
- Remove the needle and immediately dispose of it in the biohazard container.
- Apply gentle pressure to the site with a dry gauze. Do not rub or massage.
 Apply a band-aid if needed.

Intramuscular Injection Sites

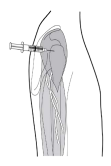


Figure 1 - Deltoid



Figure 2 - Vastus Lateralis



Figure 3 - Ventrogluteal

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Airway / Breath

Formulas

NOTE: The formulas below are for reference purposes only. Paramedics must refer to the Medical Directives and/or Base Hospital Physician patch orders for appropriate treatment options.

Cardiac/ Circula.

IV FLOW RATE CALCULATION:

qtt/min = Amount (ml) to be infused × Drops per ml (qtt/ml) of administration set Total time of infusion (min)

IOC/ Pain/ Nausea

MEDICATION INFUSION RATE:

ml/hr = Desired dose (mg/min) × 60 min/ hr

Drug concentration (mg/ml)

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Note: Units must be consistent throughout the calculation. For example, the desired dose can be in mcg/ min, as long as the concentration is also converted into mcg/ml.

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PEDIATRIC BODY WEIGHT:

For use with children aged 1 to 10 years.

(Age in years x = 2) + 10 = Approximate child body weight in kg.

Medical Refer

OXYGEN TANK DURATION:

Medic Info.

Duration of flow (minutes) = Gauge pressure - Safe residual pressure × Cylinder factor Flow rate (L/min)

Cylinder Factor: D-tank = 0.16; M-tank = 1.56

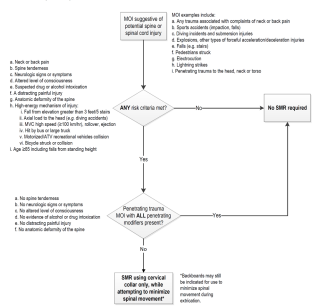
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Medical References Formulas

Spinal Motion Restriction Standard Prompt Card

This prompt card provides a quick reference of the Spinal Motion Restriction (SMR) Standard contained in the Basic Life Support Patient Care Standards (BLS PCS). Please refer to the BLS PCS for the full standard.



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"Single Strength" DOPamine Dosing Guide

DOPAMINE INFUSION RATE (mL/hr or drops/min with a microdrip set)
[Using an 800 mcg/mL ('single strength') solution]

Weight	Drip Rate (drops/min)								
(kg)	2	5	10	15	20				
		(mcg/kg/minute)							
5	1	2	4	6	8				
10	2	4	8	11	15				
15	2	6	11	17	23				
20	3	8	15	23	30				
25	4	9	19	28	38				
30	5	11	23	34	45				
35	5	13	26	39	53				
40	6	15	30	45	60				
45	7	17	34	51	68				
50	8	19	38	56	75				
55	8	21	41	62	83				
60	9	23	45	68	90				
65	10	24	49	73	98				
70	11	26	53	79	105				
75	11	28	56	84	113				
80	12	30	60	90	120				
85	13	32	64	96	128				
90	14	34	68	101	135				
95	14	36	71	107	143				
100	15	38	75	113	150				
105	16	39	79	118	158				
110	17	41	83	124	165				
115	17	43	86	129	173				
120	18	45	90	135	180				

Seizure Medical Directive Dosing Guide

Midazolam Dosing Guide

		Rou	te: IM/IN/B	uccal	R	oute: IV/I	0
Age	Weight	Supp	ose: 0.2 mg lied: 10 mg e 1 mL syri Undiluted	g/2 mL inge	Suppli Use	se: 0.1 mg ied: 10 mg 10 mL sy ed to 1 m	g/2 mL ringe
		Dose	Calculated Volume	Volume to Administer (rounded)	Dose	Actual Volume	Volume to Administer (rounded)
Neonate	3 kg	0.6 mg	0.12 mL	0.10 mL	0.3 mg	0.3 mL	0.4 mL
< 1	6 kg	1.2 mg	0.24 mL	0.25 mL	0.6 mg	0.6 mL	0.6 mL
1	12 kg	2.4 mg	0.48 mL	0.50 mL	1.2 mg	1.2 mL	1.2 mL
2	14 kg	2.8 mg	0.56 mL	0.55 mL	1.4 mg	1.4 mL	1.4 mL
3	16 kg	3.2 mg	0.64 mL	0.65 mL	1.6 mg	1.6 mL	1.6 mL
4	18 kg	3.6 mg	0.72 mL	0.70 mL	1.8 mg	1.8 mL	1.8 mL
5	20 kg	4.0 mg	0.80 mL	0.80 mL	2.0 mg	2.0 mL	2.0 mL
6	22 kg	4.4 mg	0.88 mL	0.90 mL	2.2 mg	2.2 mL	2.2 mL
			lied: 10 mg L or 10 ml			ied: 10 m 10 mL sy	
			Undiluted	١		ed to 1 m	
7	24 kg	4.8 mg	0.96 mL	1.0 mL	2.4 mg	2.4 mL	2.4 mL
8	26 kg	5.2 mg	1.04 mL	1.0 mL	2.6 mg	2.6 mL	2.6 mL
9	28 kg	5.6 mg	1.12 mL	1.2 mL	2.8 mg	2.8 mL	2.8 mL
10	30 kg	6 mg	1.20 mL	1.2 mL	3.0 mg	3.0 mL	3.0 mL
11	32 kg	6.4 mg	1.28 mL	1.2 mL	3.2 mg	3.2 mL	3.2 mL
12	34 kg	6.8 mg	1.36 mL	1.4 mL	3.4 mg	3.4 mL	3.4 mL
	40 kg	8 mg	1.60 mL	1.6 mL	4.0 mg	4.0 mL	4.0 mL
	45 kg	9 mg	1.80 mL	1.8 mL	4.5 mg	4.5 mL	4.5 mL
Max	>50 kg	10 mg	2.00 mL	2.0 mL	5.0 mg	5.0 mL	5.0 mL

Note: Dosage administered can be calculated by the weight based calculation in the Medical Directive and/or by using the above chart. Administered dosage in the chart may be rounded to the nearest volume increment that can be accurately measured.

Note:

Dosing for Adult Procedural Sedation: up tp 0.1mg/kg (IV/IM/IN); max single dose 5mg; max 2 doses

Dosing for Adult Combative Patient up to 0.1mg/kg (IV/IO/CVAD/IN); max single dose 5mg; max total dose 10mg

Medical References Seizure Medical Directcive Dosing Guide v3

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LOC/ Pain/ Nausea

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Analgesia Medical Directive - Adult & Pediatric Morphine Dosing Guide

				D-	ute: Subc					_	autas lata			
					ute: Subc ric dosag				Route: Intravenous Pediatric dosage 0.05 mg/kg					
			Supplied: 10 mg/mL					Supplied: 10 mg/mL						
									Use 1 mL Syringe					
Age	We	eight	Use 1 mL Syringe Undiluted							iluted to				
					Calcula	ted	Volum					Calculated Volume T		
			Dos	50	Volun	16	Admin (roun		Dos		Volu		Admin (round	
			Λ											
					landatory						or patier		_	
Neonate	3	kg	0.15	mg	0.015	mL		mL	0.15	mg	0.15	mL	0.15	mL
<1 1	6	kg	0.3	mg	0.03	mL	0.05	mL	0.3	mg	0.3	mL	0.3	mL
2	12	kg	0.6	mg	0.06	mL	0.05	mL	0.6	mg	0.6	mL	0.6	mL
3	14	kg	0.7	mg	0.07	mL	0.05	mL	0.7	mg	0.7	mL	0.7	mL
4	16 18	kg kg	0.8	mg	0.08	mL	0.10	mL ml	0.8	mg mg	0.8	mL ml	0.8	mL ml
5	20	kg	1.0	mg mg	0.09	mL	0.10	mL	1.0	mg	1.0	mL	1.0	mL
6	22	kg	1.1	mg	0.10	mL	0.10	ml	1.1	mg	1.0	ml	1.0	ml
7	24	kg	1.2	ma	0.11	mL	0.10	ml	1.2	mg	1.2	mL	1.2	mL
8	26	kg	1.3	ma	0.12	mL	0.1	mL	1.3	mg	1.3	mL	1.4	mL
9	28	kg	1.4	ma	0.13	mL	0.1	mL	1.4	mg	1.4	mL	1.4	mL
10	30	kg	1.5	mg	0.14	mL mL	0.1	mL	1.5	ma	1.5	mL	1.6	mL
11	32	kg	1.6	ma	0.15	mL mL	0.2	mL	1.6	ma	1.6	mL	1.6	mL
				J	0.10	IIIL								
					ıpplied: 1 Jse 1 mL :						pplied: 1 se 10 mL			
					Undilu		,•				iluted to			
	34	kg	1.7	mg	0.17	_	0.2	ml	1.7	mg	1.7	ml	1.8	ml
	40	kg	2.0	mg	0.17	mL	0.2	ml	2.0	mg	2.0	ml	2.0	ml
	45	kg	2.25	mg	0.225	mL mL	0.2	mL	2.25	mg	2.25	mL	2.2	mL
	50	kg	2.5	mg	0.25	mL	0.3	mL	2.5	mg	2.5	ml	2.6	mL
	55	kg	2.75	mg	0.25	mL	0.3	mL	2.75	mg	2.75	mL	2.8	mL
	60	kg	3.0	mg	0.275	mL	0.3	mL	3.0	mg	3.0	mL	3.0	mL
	65	kg	3.25	mg	0.325	mL	0.3	mL	3.25	mg	3.25	mL	3.2	mL
Youth	70	kg	3.5	mg	0.325	mL	0.4	mL	3.5	mg	3.5	mL	3.6	mL
(12-17)	75	kg	3.75	mg	0.375	mL	0.4	mL	3.75	mg	3.75	mL	3.8	mL
	80	kg	4.0	ma	0.375		0.4	mL	4.0	ma	4.0	mL	4.0	mL
	85	Kg	4.25	ma	0.40	mL	0.4	mL	4.25	mg	4.25	mL	4.2	mL
	90	kg	4.5	mg		mL	0.5	mL	4.5	mg	4.5	ml	4.6	mL
	95	kg	4.75	mg	0.45	mL	0.5	mL	4.75	mg	4.75	mL	4.8	mL
	100	kg	4.75	mg	0.475	mL	0.5	mL	5.0	mg	5.0	mL	5.0	mL
Pediatrio	: Maximu			-	0.5	mL								
	Dose				0.50				5.0					

Dosing Interval: 15 minutes Pediatric Max # of Doses: 4

Analgesia Medical Directive - Adult & Pediatric Morphine Dosing Guide

	Use 1 m	10 mg/mL L Syringe lluted	Supplied: Use 10 ml Diluted to	_ Syringe
Adult N/A	2 - 10mg	0.2 - 1.0 mL	2 – 10 mg	2 - 10 mL
Adult Maximum Single Dose	10 mg	1.0 mL	10 mg	10 mL

Dosing Interval: 15 minutes Adult Max # of Doses: 4

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Analgesia Medical Directive - Adult & Pediatric FentaNYL Dosing Guide

Route: Intravenous or Intranasal Supplied: 100 mcg in 2 mL *Intranasal Max Fluid : 1 mL per nare Use 1 mL Syringe, undiluted

Maximum Pediatric Dosage: up to 1 mcg/kg (administer in divided doses)

⚠ Mandatory Provincial Patch Point for Children < 12 years old					
Age	Weight	Maximum Dose	Calculated Volume	Volume to administer (rounded)	
Neonate	3 kg	3 mcg	0.03 mL		
<1	6 kg	6 mcg	0.06 mL	0.05 mL	
1	12 kg	12 mcg	0.24 mL	0.2 mL	
2	14 kg	14 mcg	0.28 mL	0.3 mL	
3	16 kg	16 mcg	0.32 mL	0.3 mL	
4	18 kg	18 mcg	0.36 mL	0.4 mL	
5	20 kg	20 mcg	0.40 mL	0.4 mL	
6	22 kg	22 mcg	0.44 mL	0.4 mL	
7	24 kg	24 mcg	0.48 mL	0.5 mL	
8	26 kg	26 mcg	0.52 mL	0.5 mL	
9	28 kg	28 mcg	0.56 mL	0.6 mL	
10	30 kg	30 mcg	0.60 mL	0.6 mL	
11	32 kg	32 mcg	0.64 mL	0.6 mL	
	34 kg	34 mcg	0.68 mL	0.7 mL	
	40 kg	40 mcg	0.80 mL	0.8 mL	
	45 kg	45 mcg	0.90 mL	0.9 mL	
Youth*	50 kg	50 mcg	1.0 mL	1.0 mL	
(12-17)	55 kg	55 mcg	1.1 mL*	1.1 mL*	
	60 kg	60 mcg	1.2 mL*	1.2 mL*	
	65 kg	65 mcg	1.3 mL*	1.3 mL*	
	70 kg	70 mcg	1.4 mL*	1.4 mL*	
	75 kg	75 mcg	1.5 mL*	1.5 mL*	
	Maximum e Dose*	75 mcg	1.5 mL*	1.5 mL*	
Adults ≥	≥ 18 years	25 – 75 mcg	0.50 -1.5 mL*	0.50 -1.5 mL*	
	imum Single ose	75 mcg	1.5 mL*	1.5 mL*	

^{*}for pediatric dosing, consider administering in divided doses of one-third to one-half and titrate to effect similar to adult dosing.

EPINEPHrine 1 mg/mL = 1:1000 IM Dosing Guide

Dose (0.01 mg/kg) is rounded to the nearest 0.05mg Use a 1 mL syringe

AGE	WEIGHT	DOSE (mg)	VOLUME (mL) (rounded)
3 months	5 kg	0.05 mg	0.05 mL
6 months	8 kg	0.08 mg	0.10 mL
9 months	10 kg	0.10 mg	0.10 mL
1 year	12 kg	0.12 mg	0.10 mL
2 years	14 kg	0.14 mg	0.15 mL
3 years	16 kg	0.16 mg	0.15 mL
4 years	18 kg	0.18 mg	0.20 mL
5 years	20 kg	0.10 mg	0.20 mL
,	ŭ		
6 years	22 kg	0.22 mg	0.20 mL
7 years	24 kg	0.24 mg	0.25 mL
8 years	26 kg	0.26 mg	0.25 mL
9 years	28 kg	0.28 mg	0.30 mL
10 years	30 kg	0.30 mg	0.30 mL
11 years	32 kg	0.32 mg	0.30 mL
12 years	34 kg	0.34 mg	0.35 mL
13 years	36 kg	0.36 mg	0.35 mL
14 years	38 kg	0.38 mg	0.40 mL
Adult	50 kg	0.50 mg	0.50 mL

Note: Dosage administered can be calculated by the weight based calculation in the Medical Directive and/or by using the above chart. Administered dosage in the chart may be rounded to the nearest volume increment that can be accurately measured. 111110

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Medication Information

PRIMARY CARE PARAMEDIC MEDICAL DIRECTIVES

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Medication Information

	ACETAMINOPHEN
CLASS	Antipyretic and analgesic. Mild anti-inflammatory effects.
ACTION	Exact mechanism is not known. Rapidly absorbed through GI tract. Believed to raise the pain threshold.
ONSET	15 minutes and lasts up to 3 hours.
METABOLISM	At normal therapeutic dosages, primary hepatic metabolism. A toxic dose (as little as 4g daily) can cause hepatic cell necrosis. Oral administration is subject to first pass metabolism.
	ADENOSINE
CLASS	Antiarrhythmic
ACTION	Slows conduction time through the AV node, interrupting the re-entry pathways through the AV node, restoring normal sinus rhythm. Adenosine also causes coronary vasodilation and increases blood flow in normal coronary arteries with little to no increase in stenotic coronary arteries; thallium-201 uptake into the stenotic coronary arteries will be less than that of normal coronary arteries revealing areas of insufficient blood flow.
ONSET	Rapid
HALF-LIFE	< 10 seconds
METABOLISM	Blood and tissue.
	AMIODARONE
CLASS:	Antiarrhythmic (Class I, II, III, and IV)
ACTION:	Blocks sodium channels; lengthens cardiac potential. Slows cardiac conduction through the AV node. Antisympathetic action and negative inotropic effects in cardiac nodal tissue. Used for ventricular arrhythmias (ventricular tachycardia/ventricular fibrillation) and some atrial arrhythmias (atrial fibrillation, but takes hours)
ONSET	15 minutes
TIME TO PEAK	1 to 4 hours
DURATION	3 to 6 hours
HALF-LIFE	9-36 hours
METABOLISM	Hepatic

Inti	റ

	ASPIRIN (ASA)
CLASS:	Platelet aggregation inhibitor, analgesic, antipyretic and anti- inflammatory
ACTION:	Decreases clotting by inactivating cyclooxygenase, interfering with Thromboxane A2 production within the platelets. Thromboxane A2 also causes arteries to constrict. Reduced morbidity/mortality in adults with C/P from an AMI.
ABSORPTION	Rapid
TIME TO PEAK	1-2 hours
METABOLISM	Hydrolyzed to salicylate (active) in GI mucosa, RBC, synovial fluid and blood. Metabolism of salicylate primarily by the liver.

	ATROPINE
CLASS	Parasympatholytic, anticholinergic
ACTION	Blocks the action of acetylcholine at parasympathetic sites in smooth muscle, secretory glands and the CNS. Results in increased cardiac output and dries secretions.
ONSET	Rapid
HALF-LIFE	2-3 hours
DISTRIBUTION	Widely throughout the body; crosses placenta; trace amounts enter breast milk; crosses blood-brain barrier.
METABOLISM	Hepatic

	CALCIUM GLUCONATE 10%
CLASS	Minerals and electrolytes
ACTION	Calcium protects the myocardium from the deleterious effects of hyperkalemia. It stabilizes the cardiac cell membrane.
ADVERSE REACTION	When given too rapidly can cause hypotension, bradycardia and syncope. If administered IM or extravagates it can cause necrosis/abscess. When given to someone on digoxin this may cause sudden death from ventricular fibrillation.
ADMIN	Slow IV push over 2-3 minutes Incompatible with Sodium Bicarbonate in same IV line.
ONSET	Rapid
DURATION	30 minutes - 2hours
SIDE EFFECTS	Chalky taste, N&V, Dry mouth

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DEXAMETHASONE CLASS Adrenocoritcal steroid Binds to the glucocorticoid receptors inhibiting the release ACTION of pro-inflammatory signals through cytokine inhibition. resulting in decreased edema, fibrin deposition, capillary leakage and migration of inflammatory cells. ONSET 5-15 min(IV); 30 min (PO)60 minutes DURATION 3 days HALF-LIFE

4 hours

Cardiac/ Circula.

> DEXTROSE (D50) IN WATER CLASS Carbohydrate ACTION Replenishes blood glucose levels, reversing hypoglycemia. METABOLISM Metabolized to carbon dioxide and water.

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	DIMENHYDRINATE (GRAVOL)
CLASS	Antiemetic, Antihistamine
ACTION	Competes with histamine for H1-receptor sites on effector cells in the GI tract, blood vessels and respiratory tract; blocks chemoreceptor trigger zone, diminishes vestibular stimulation and depresses function through its central anticholinergic activity.
ONSET	1-5 minutes (IV). 15-30 minutes (oral)
PEAK EFFECTS	1-2 hours
DURATION	3-6 hours

DIPENHYDRAMINE (BENADRYL) CLASS Antihistamine ACTION Competes with histamine and H1-receptor sites on effector cells in the GI tract, blood vessels and respiratory tract; anticholinergic and sedative effects are also seen. ONSET 1-5 minutes (IV), 1-3 hours (oral) 1-2 hours (IV). 2-4 hours (oral) PEAK FEFECTS HALF-LIFE 2-10 hours DURATION 4-8 hours

DOPAMINE
Sympathomimetic agent
Stimulates both adrenergic and dopaminergic receptors, lower doses are mainly dopaminergic stimulating and produce renal and mesenteric vasodilation. Higher doses have both dopaminergic and \$1-adrenergic stimulating and produce cardiac stimulation and renal vasodilation. Large doses stimulate d-adrenergic receptors.
5 minutes
2 minutes
Renal, hepatic and plasma (25% gets converted to norepinephrine).

	EPINEPHERINE
CLASS	Sympathomimetic agent
ACTION	Stimulate $\beta 1$, $\alpha 1$ and $\beta 2$ -adrenergic receptors resulting in relaxation of smooth muscle of the bronchial tree, cardiac stimulation (increasing myocardial O2 consumption) and dilation of skeletal muscle vasculature. Small doses can cause vasodilation via $\beta 2$ -vascular receptors; large doses may produce constriction of skeletal and vascular smooth muscle.
ONSET	5-10 minutes (bronchodilation).
METABOLISM	Hepatic

	FENTANYL
CLASS	Analgesic, opioid
ACTION	Binds to opioid mu-receptors in the CNS causing inhibition of ascending pain pathways, altering the perception of and response to pain; produces generalized CNS depression, respiratory depression, and can cause apnea. Can cause muscle rigidity if rapid IV injection.
ONSET	IV: almost immediately IN: 5-15 minutes
PEAK EFFECT	IV: 6 minutes IN: 12 minutes
METABOLISM	Hepatic

Intro

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic.

Contact

GLUCAGON CLASS Glucose elevating agent ACTION Stimulates adenylate cyclase to produce increased cyclic AMP, which promotes hepatic glycolysis and gluconeogenesis. ONSET 30 minutes (IM)

Cardiac/ Circula.

resulting in a rise in blood glucose levels. HALF-LIFE 8-18 minutes DURATION 60-90 minutes METABOLISM Primarily hepatic, some occurs renally and in the plasma.

IOC/ Pain/ Nausea

GLYCOPYRROLATE CLASS anticholinergic

Inhibits the acetylcholine activity on smooth muscles and structures innervated by postganglionic nerves. Causes bronchodilation, decreased volume and acidity of gastric

secretions, as well as control of excessive pharvngeal, tracheal and bronchial secretions. Also has antimuscarinic properties, antagonizes muscarinic effects induced by cholinergic medications

Possible effect through central dopamine, adrenergic, cholinergic

Proced.

ONSET Rapid DURATION 2-4 hours HALF-LIFE 1.25 hours

Research/ Sp. Proj

HALOPERIDOL

Medical Refer

CLASS Antipsychotic ACTION Butyrophenone antipsychotic unclear mechanism of action.

and histaminergic receptors. ONSET Rapid

DURATION

ACTION

4-6 hours

Contact

Destinat Guide.

Medication Information

HYDROCORTISONE		
CLASS	Adrenal glucocorticoid, corticosteroid	
ACTION	Short-acting corticosteroid; when used in adrenal crisis or adrenocortical deficiency it replaces/mimics the person's own cortisol which regulates glucose, regulates the immune system, and is released during stressors to help support the cardiovascular system	
ONSET	1-2 hours	
PEAK EFFECT	1.5 – 2 hours	
DURATION	6-12 hours	
METABOLISM	Hepatic	

	HYDROMORPHONE
CLASS	Opioid analgesic
ACTION	Binds to the mu-opioid receptors in the CNS causing inhibition of the ascending pain pathways, altering the perception of and response to pain. Produces generalized CNS depression
ONSET	5 minutes
DURATION	3-4 hours
HALF-LIFE	2-3 hours

	IBUPROFEN
CLASS	Antipyretic, analgesia and non-steroid anti-inflammatory
ACTION	Its pharmacological effects are believed to be due to inhibition COX-2 which decreases the synthesis of prostaglandins involved in mediating inflammation, pain, fever and swelling. Antipyretic effects may be due to action on the hypothalamus, resulting in an increased peripheral blood flow, vasodilation, and subsequent heat dissipation.
PEAK EFFECT	120 minutes
ONSET	15 minutes
DURATION	4-6 hours
ADVERSE EFFECTS	HTN, MI, GI bleeding, increased the risk of gastric ulcers and damage and renal failure.
METABOLISM	Ibuprofen and its metabolites pass easily across the placenta. More than 90% of an ingested dose is excreted in the urine as metabolites or their conjugates.

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medio

Contact

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer.

Medio

Contact

	·
	KETOROLAC (TORADOL)
CLASS	Analgesic, antipyretic and non-steroid anti-inflammatory
ACTION	Blocks prostaglandin formation thereby decreasing nociceptor stimulation.
ONSET	10 minutes (IM/IV)
PEAK EFFECT	2-3 hours
DURATION	6-8 hours
METABOLISM	Mostly the hepatic

LIDOCAINE (XYLOCAINE)	
CLASS	Class 1b antiarrhythmic
ACTION	Suppresses automaticity of conductive tissue by increasing the electrical stimulus threshold of the ventricles, His-Purkinje system and spontaneously depolarization of the ventricles during diastole (by direct action on the tissues). Blocks both the initiation and conduction of nerve impulses by decreasing the neural membranes permeability to Na ions, which results in inhibition of depolarization with resultant blockade of conduction.
ONSET	45-90 seconds
DURATION	10-20 minutes
METABOLISM	90% hepatic

	MIDAZOLAM (VERSED)
CLASS	Benzodiazepine, CNS depressant, Sedative and Amnesic
ACTION	Binds to stereospecific benzodiazepine receptors on the post- synaptic GABA neuron at several sites within the CNS (including limbic system and reticular formation). Enhancement of the inhibitory effect of GABA on neural excitability results by increased neural membrane permeability to chloride ions. This shift in chloride.
ONSET	45-90 seconds
DURATION	10-20 minutes
METABOLISM	90% hepatic

	MORPHINE
CLASS	Opioid analgesia
ACTION	Binds to opiate receptors in the CNS causing inhibition of ascending pain pathways, altering the perception of and response to pain; produces generalized CNS depression.
ONSET	2-5 minutes (IV)
PEAK EFFECT	20 minutes (IV)
METABOLISM	Hepatic

NALOXONE (NARCAN)		
CLASS	Narcotic Antagonist	
ACTION	Competitive narcotic antagonist. Displaces and narcotics bound to opiate receptor sites reversing their effects.	
ONSET	2-5 minutes (IM). 8-18 minutes (IN). 2 minutes (IV)	
HALF-LIFE	3-4 hours (neonates). 0.5-1.5 hours (adults)	
DURATION	30-120 minutes	
DISTRIBUTION	Crosses placenta	
METABOLISM	Hepatic	

	NITROGLYCERIN
CLASS	Coronary vasodilator, smooth muscle relaxant and anti-anginal
ACTION	Vasodilation of peripheral veins and arteries with more prominent effects on the veins. Reduces myocardial oxygen demand by decreasing preload; may modestly reduce afterload; dilates coronary arteries and improves collateral flow to ischemic tissues. In smooth muscle, nitric oxide activates smooth muscle relaxation.
ONSET	1-3 minutes (SL). 15-30 minutes (topical). 30 minutes (transdermal)
HALF-LIFE	1-4 minutes
DURATION	25 minutes (SL), 7 hours (topical), 10-12 hours (transdermal)
METABOLISM	Extensive first-pass effect; hepatic, RBC and vascular walls

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer.

Medic

Contact

ONDANSETRON	
CLASS	5-HT3 antagonist
ACTION	Selective 5-HT3 receptor antagonist. Mechanism of action through blocking the action of 5-HT3 selectively peripherally and through the vagus nerve, a natural substance that may cause nausea and vomiting. Centrally the chemoreceptor trigger zone is effected.
ONSET	20-30 min
HALF-LIFE	3-6 hrs (PO); 5-8 HRS (IV, IM)
DURATION	4-8 hrs (PO); 5-8 hrs (IV, IM)

OXYTOCIN		
CLASS	Hormone	
ACTION	Promotes uterine contractions by increasing intracellular calcium levels. Greatest effect during labor at term due to increased oxytocin receptor concentrations in uterine myometrial tissue	
ONSET	3-5 min	
HALF-LIFE	2-3 hrs	
DURATION	1-6 min	

	SALBUTAMOL (VENTOLIN)
CLASS	Sympathomimetic, β2 agonist
ACTION	Relaxes bronchial smooth muscle by action on $\beta 2\mbox{-receptors}$ with little effect on heart rate
ONSET	10 minutes (Neb/Inhalation)
HALF-LIFE	3-8 hours (inhaled)
DURATION	3-4 hours (Neb/Inhalation)
METABOLISM	Hepatic to an inactive sulfate

	XYLOMETAZOLINE (OTRIVIN)
CLASS	Sympathomimetic Adrenergic Alpha-agonist, decongestant
ACTION	When sprayed into the nares, causes vasoconstriction of the nasal mucosa, resulting in a decrease in blood flow in the nasal passages, decreased nasal congestion, and may help stop epistaxis.
ONSET	5-10 minutes

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic.

Contact

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic Info.

Contact

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Medication Information



For the Paramedic:

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic.

If a paramedic encounters a physician on-scene that would like to assist or direct care, the paramedic will follow the Ontario BLS-PCS for any BLS care and the Medical Directives in this document for any ALS care. Inform the physician that paramedics are not able to accept delegation for controlled medical acts from any physician other than an affiliated Base Hospital Physician. The paramedic may show the following information to the on-scene physician to assist in next steps and provide further information.

Physician On-Scene Reference

To the On-Scene Physician:

Thank you for your assistance.

The paramedics would usually take responsibility for the patient(s) upon their arrival. If, as a physician, you wish to assist with the emergency after the providers have arrived you have three options:

- 1. Offer your assistance or suggestions that follow the Ontario Basic Life Support Patient Care Standards and/or the Paramedic Medical Directives. If your instructions are not in accordance with these documents, the paramedics cannot follow this direction but can contact the Regional Base Hospital Physician for direction.
- 2. Take complete responsibility for patient in which case you will need to accompany the patient to hospital. The paramedics will assist you, but cannot perform skills that do not follow their directives. You may be asked to show identification that you are a physician licensed to practice medicine in Ontario.
- Request to speak with the Regional Base Hospital Physician (via patch) to offer advice and consult on the best management of the patient(s).

Contact



Enhanced dedicated BHP support for Paramedic consultation and new Patient Care Models

CPER REGIONAL PATCH NUMBER 1-888-554-8011



identification

Identify BHP Introduce yourself

S SITUATION

ORDERS SOUGHT age, sex, weight problem / concern ETA to hospital

B BACKGROUND

Pertinent +/-HPI (OPORST) PMHx (SAMPLE)

ASSESSMENT

Pertinent +/-Physical Exam Vitals Signs, ECG

R RESPONSE

Response to treatment Reiterate orders sought Receive orders REPEAT BACK ORDERS



CONNECTION ISSUES

Use CACC // Radio Backup to reach OMC BHP



Hamilton Health Sciences



Intro

Airway / Breath.

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Contact Information

430 McNeilly Road, Unit 201 Stoney Creek, Ontario L8E 5E3

Telephone Number: 905-521-2100 x71223 Fax Number: 905-643-1104

Position:	EXT:	Mobile:	Email Address:
Regional Program Manager/ Director		905-515-4818	tdodd@cper.ca
Regional Medical Director			millerpa@hhsc.ca
Assistant Medical Director			rupinder@sahsi.net
Assistant Medical Director			erich.hanel@medportal.ca
Senior Medical Advisor			agarg@mcmaster.ca
Administration Assistant (To the Directors)	71226		ceaston@cper.ca
Administration Assistant (To the Programs)	71229		acollie@cper.ca
Administration Assistant	71223		jswing@cper.ca
Lead Quality Specialist		289-286-0975	aburgess@cper.ca
Quality Specialist		905-870-4457	kselvar@cper.ca
Quality Specialist		519-503-6632	cschneider@cper.ca
Quality Specialist		416-436-5428	winterkat@hhsc.ca
Lead Paramedic Educator		905-515-0659	scoletta@cper.ca
Paramedic Educator		289-219-1952	dplyley@cper.ca
Paramedic Educator		289-260-3268	jradoslav@cper.ca
Paramedic Educator			kapadiab@hhsc.ca
Outreach Specalist		365-324-8389	pdeath@cper.ca
	Regional Program Manager/ Director Regional Medical Director Assistant Medical Director Assistant Medical Director Senior Medical Advisor Administration Assistant (To the Directors) Administration Assistant (To the Programs) Administration Assistant (To the Programs) Administration Assistant Lead Quality Specialist Quality Specialist Quality Specialist Quality Specialist Lead Paramedic Educator Paramedic Educator Paramedic Educator	Regional Program Manager/ Director Regional Medical Director Assistant Medical Director Assistant Medical Director Senior Medical Advisor Administration Assistant (To the Directors) Administration Assistant (To the Programs) Administration Assistant (To the Programs) Administration Assistant (To the Programs) Administration Assistant Quality Specialist Quality Specialist Quality Specialist Quality Specialist Lead Paramedic Educator Paramedic Educator Paramedic Educator	Regional Program Manager/ Director 905-515-4818 Regional Medical Director Assistant Medical Director Assistant Medical Director Assistant Medical Director Senior Medical Advisor Administration Assistant (To the Directors) 71226 Administration Assistant (To the Programs) 71229 Administration Assistant (To the Programs) 71223 Lead Quality Specialist 289-286-0975 Quality Specialist 905-870-4457 Quality Specialist 519-503-6632 Quality Specialist 416-436-5428 Lead Paramedic Educator 905-515-0659 Paramedic Educator 289-219-1952 Paramedic Educator 289-260-3268 Paramedic Educator 289-260-3268

Cardiac/ Circula.

Airway /

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Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

HHS Centre for Paramedic Education and Research Additional Contact **Information Reference**

Central Ambulance Communication Centres (CACC):

800-265-2215 CACC - Cambridge CACC - Hamilton 905-574-1414 CACC - Hamilton (Alternate) 800-263-5767 CACC - Niagara Ambulance Communication Centre 905-704-4005 866-895-6227

Emergency Medical Services:

Brant / Brantford Paramedic Service 519-756-4570 **Dufferin County Paramedic Service** 519-941-9608 Guelph-Wellington Paramedic Service 519-824-1677 Haldimand County Paramedic Services 905-318-5932 Hamilton Paramedic Service 905-546-2424 905-641-0827 Niagara EMS Norfolk County Paramedic Services 519-426-4115 Region of Waterloo Paramedic Service 519-650-8295 Six Nations Paramedic Services 519-445-4000

Airway / Breath

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced

Research / Sp. Proj

Medical Refer.

Medic. Info

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

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Medic. Info.

Contact

Community Support Referral Contact Information

The following contact information is provided for cases where:

- Patients are refusing transport to the hospital, and
- An assessment shows that the patient has the capacity to refuse, and
- The patient does not appear to be of immediate danger to themselves or others, and
- Paramedics have ongoing concerns regarding the living conditions in their home (CCAC), their need for victim's support services (victim's services) or the patient's mental health (COAST, Hamilton only)
- ▶ OR the family of a patient needs support services (Victims Services).

These community service organizations are available to assist people with these concerns. Paramedics can give the information directly to the patient or assist them by making the referral on their behalf. Please note that if the Paramedic assists the patient by calling the organization he/she must_get the patient's consent to do so. If the Paramedic contacts the organization directly, the agency will require the patient's name, address, phone number and nature of the concern. The Paramedic must then leave the information about the organization called with the patient.



CCAC (Community Care Access Centre): provides services for persons with living condition concerns (message can be left).

 Brantford CCAC:
 800-810-0000

 Dufferin County CCAC:
 519-925-5452

 Guelph-Wellington CCAC:
 519-823-2550

 Haldimand / Hamilton CCAC:
 800-810-0000

 Niagara Region CCAC:
 800-810-0000

 Norfolk / Simcoe CCAC:
 800-810-0000

 Six Nations (Ohsweken)
 519-445-2418

 Waterloo - Kitchener CCAC:
 519-748-2222

Victims Services: provides short-term emotional support and community referral and assistance to victims of crime, tragic circumstance or disaster (24/7).

Brantford	519-752-3140
Cambridge	519-585-2369 / 519-570-5143
Dufferin County	519-942-1452
Guelph-Wellington	519-824-1212 ext. 7304
Haldimand County	800-264-6671
Hamilton Victim Services	905-546-4904
Kitchener	519-585-2369 / 519-570-5143
Niagara Region	905-682-2626
Norfolk County	800-264-6671
Six Nations (Ohsweken)	519-752-3140
Waterloo Region	519-585-2369 /



COAST (Crisis Outreach And Support Team): provides services for persons with mental health concerns in the Hamilton area only (24/7).

519-570-5143

Hamilton - Only (24/7) 905 972-8338 Intro

Airway / Breath.

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced

Research/ Sp. Proj

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Proced.

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Medic.

Contact

Child in Need of Protection

Paramedics have a duty to report under the Child and Family Services Act (CFSA) and this extends to any child they encounter in their professional duties and is not limited to the person (s) requesting 9-1-1 services¹. This duty overrides any other provincial statue, including any provisions that would otherwise prohibit someone from making a disclosure (i.e. PHIPA). This failure to report a suspicion in the circumstances set out in the CFSA is an offence under the Act.²

Children's Aid Societies in Ontario

Dufferin Child and Family Protection Services

...,

Bus: (519) 941-1530

Family & Children's Services of Guelph and Wellington County

v.

Bus: (519) 824-2410

Children's Aid Society of Hamilton

Bus: (905) 522-1121

Catholic Children's Aid Society of Hamilton Bus: (905) 525-2012

Family & Children's Services Niagara Bus: (888) 937-7731

Children's Aid Society of Haldimand and Norfolk Bus: (519) 587-5437 Toll Free: (888) 227-5437

Brant Family and Children's Services Bus: (519) 753-8681 Toll Free: (888) 753-8681

Family & Children's Services of the Waterloo Region

Bus: (519) 576-0540

¹Training Bulletin 116 -Child in Need of Protection Standard March 2015 Version 1.0

² Basic Life Support Patient Care Standards –Version 2.2

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Airway / Breath.

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Proced.

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Airway / Breath.

Cardiac/ Circula.

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Destination Guidelines

DRIMARY CARE BARAMERIC MEDICAL DIRECTIVES

Cardiac/ Circula.

IOC/

Pain/

Nausea

Proced

Research / Sp. Proj

Medical

Refer.

Field Trauma Triage Standards

Definitions

For the purposes of the Field Trauma Triage Standard:

Regionally Designated Equivalent Hospital

means an appropriately resourced hospital facility as defined by the Regional Trauma Network of Critical Care Services Ontario and included in a local PPS.

Transport Time

means the time from scene departure to time of arrival at destination.

General Directive

The paramedic shall follow the procedure below when conducting field triage of patients injured by a traumatic mechanism or who show evidence of trauma.

The paramedic shall also use this standard to assess the clinical criteria (i.e. to determine if the patient meets the clinical criteria) as required by the Air Ambulance Utilization Standard.

The paramedic shall consider using the Trauma Termination of Resuscitation (TOR) contained in the Trauma Cardiac Arrest Medical Directive as per the ALS PCS.

CACC/ACS may authorize the transport once notified of the patient's need for re-direct or transport under the Field Trauma Triage Standard.

Procedure

The paramedic shall:

- 1. assess the patient to determine if he/she has one or more of the following physiological criteria (Step 1):
 - Patient does not follow commands.
 - b. Systolic blood pressure <90mmHg, or
 - c. Respiratory rate <10 or ≥30 breaths per minute or need for ventilatory support (<20 in infant aged <1 year);
- if the patient meets the physiological criteria listed in paragraph 1 above, AND the land 2. transport time is estimated to be <30 minutes* to a Lead Trauma Hospital (LTH) or regionally designated equivalent hospital, transport the patient directly to the LTH or regionally designated equivalent hospital;
- 3. if the patient does not meet the criteria listed in paragraphs 1 and 2, assess the patient to determine if he/she has one or more of the following anatomical criteria (Step 2):

Medic. Info.

- a. Any penetrating injuries to head, neck, torso and extremities proximal to elbow or knee.
- b. Chest wall instability or deformity (e.g. flail chest),
- c. Two or more proximal long-bone fractures,
- d. Crushed, de-gloved, mangled or pulseless extremity,
- e. Amputation proximal to wrist or ankle,
- f. Pelvic fractures.
- g. Open or depressed skull fracture, or
- h. Paralysis;
- 4 if the patient meets the anatomical criteria listed in paragraph 3 above and the land transport time is estimated to be <30 minutes* to the LTH or regionally designated equivalent hospital, transport the patient directly to the LTH or regionally designated equivalent hospital:
- 5 if unable to secure the patient's airway or survival to the LTH or regionally designated equivalent hospital is unlikely, transport the patient to the closest emergency department despite paragraphs 2 and 4 above;
- 6 despite paragraph 5 above, transport the patient directly to an LTH or regionally designated equivalent hospital if the patient has a penetrating trauma to the torso or head/neck, and meets ALL of the following:
 - a. Vital signs absent vet not subject to TOR described in the General Directive above.
 - b. Land transport to the LTH or regionally designated equivalent hospital is estimated to be <30 minutes*:
- 7. if the patient does not meet the physiological or anatomical criteria listed above, use the following criteria to determine if the patient may require other support services at the LTH or regionally designated equivalent hospital as a result of his/her traumatic mechanism of injury (Step 3):
 - a. Falls
 - i. Adults: falls ≥6 metres (one story is equal to 3 metres)
 - ii. Children (age <15): falls >3 metres or two to three times the height of the child
 - b. High Risk Auto Crash
 - i. Intrusion ≥0.3 metres occupant site; ≥0.5 metres any site, including the roof
 - ii. Ejection (partial or complete) from automobile
 - iii. Death in the same passenger compartment
 - iv. Vehicle telemetry data consistent with high risk injury (if available)
 - c. Pedestrian or bicyclist thrown, run over or struck with significant impact (≥30 km/hr) by an automobile
 - d. Motorcycle crash ≥30 km/hr;
- 8. if the patient meets the mechanism of injury criteria listed in paragraph 7 above, AND the land transport time is estimated to be <30 minutes* to an LTH or regionally designated equivalent hospital, determine the need for patient transport to the LTH or regionally designated equivalent hospital;

Airway / Breath

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced

Research / Sp. Proj

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Medic. Info

Airway / Breath.

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Proced.

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Medic Info.

Contact

in conjunction with the physiological, anatomical, and mechanism of injury criteria listed above, consider the following special criteria (Step 4):

a. Age i. Risk of injury/death increases after age 55

ii. SBP <110 may represent shock after age 65

b. Anticoagulation and bleeding disorders

c. Burns

9.

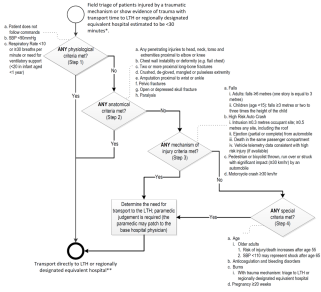
i. With trauma mechanism: triage to LTH

d. Pregnancy ≥20 weeks; and

10. if the patient meets any of the special criteria listed above, AND the land transport time is estimated to be <30 minutes* to an LTH or regionally designated equivalent hospital, determine the need for patient transport to the LTH or regionally designated equivalent hospital.

*Note: The 30 minute transport time may be amended to up to 60 minutes as per an ambulance service PPS, but may not exceed 60 minutes.

This prompt card provides a quick reference of the Field Trauma Triage Standard contained in the Basic Life Support Patient Care Standards (BLS PCS). Please refer to the BLS PCS for the full standard.



^{*}The 30 minute transport time may be amended to up to 60 minutes as per an ambulance service PPS, but may not exceed 60 minutes

^{**}If unable to secure the patient's airway or survival to the LTH or regionally designated equivalent hospital is unlikely, transport the patient to the closest ED (unless patient has penetrating trauma to the torso or head/neck). Consider the Trauma TOR as per the ALS PCS.



Airway / Breath.

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced

Research/ Sp. Proj

Medical Refer.

Medic. Info

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer.

Medic.

Contact

Air Ambulance Utilization Standard

General Directive

Requests for an on-scene air ambulance response should meet at least one of the bulleted operational criteria **PLUS** one of the clinical criteria (e.g. known clinical criteria as listed in the *Field Trauma Triage Standard* or from the bulleted list of medical or obstetrical criteria listed below).

Procedure

The paramedic shall:

- 1. assess the scene response to meet one or more of the following operational criteria:
 - a. The land ambulance is estimated to require more than 30 minutes to reach the scene and the air ambulance can reach the scene quicker.
 - b. The land ambulance is estimated to require more than 30 minutes to travel from the scene to the closest appropriate hospital* and the air ambulance helicopter can reach the scene and transport the patient to the closest appropriate hospital* quicker than the land ambulance.
 - c. The estimated response for both land and air is estimated to be greater than 30 minutes, but approximately equal, and the patient needs care which cannot be provided by the responding land ambulance.
 - d. There are multiple patients who meet the clinical criteria and the local land ambulance resources are already being fully utilized.
- if the scene response meets the requirements of paragraph 1 above, assess the patient to determine if he/she meets one or more of the following clinical criteria:
 - a. Patients meeting the criteria listed in the Field Trauma Triage Standard.
 - b. Patients meeting one or more of the following:

i. Medical:

- Shock, especially hypotension with altered mentation (e.g. suspected aortic aneurysm rupture, massive gastrointestinal bleed, severe sepsis, anaphylaxis, cardiogenic shock, etc.)
- 2. Acute stroke with a clearly determined time of onset or last known to be normal <6.0 hours
- 3. Altered level of consciousness (GCS <10)
- 4. Acute respiratory failure or distress
- 5. Suspected STEMI or potentially lethal dysrhythmia
- 6. Resuscitation from respiratory or cardiac arrest
- 7. Status epilepticus
- 8. Unstable airway or partial airway obstruction

ii. Obstetrical:

- Active labour with abnormal presentation (i.e. shoulder, breech or limb)
- 2. Multiple gestation and active labour
- 3. Umbilical cord prolapse
- Significant vaginal bleeding (suspected placental abruption or placenta previa or ectopic pregnancy);
- in conjunction with the ACO, assess if an on-scene air ambulance helicopter is appropriate, based on:
 - a. the perceived severity of the reported injuries and without confirmation that the clinical criteria have been met, or
 - the patient cannot reasonably be reached by land ambulance (e.g. sites without road access such as islands; geographically isolated places, etc.);
- 4. if the requirements listed in paragraph 2 or 3 above are met, request an on-scene air ambulance helicopter response:
 - a. Provide the ACO with the information set out in operational and clinical criteria above. In order for the ACO to determine if an air ambulance response and transport will be quicker than land ambulance, the paramedic will provide the ACO with the estimated time to prepare the patient for transport, identify separately any time required for patient extrication, provide the estimated land ambulance driving time to the closest appropriate hospital and any additional information as required.
 - b. The paramedics shall not delay patient transport by waiting for the air ambulance helicopter, unless the air ambulance helicopter can be seen on its final approach to the scene. If the air ambulance helicopter is en route but not on final approach to the scene, and the land paramedics have the patient in his/her ambulance, then the land ambulance will proceed to the closest local hospital with an emergency department. The air ambulance helicopter will proceed to that local hospital and, if appropriate, assist hospital personnel prepare the patient for rapid evacuation.
 - while en route to the local hospital, paramedics may rendezvous with the air ambulance helicopter if:
 - the air ambulance helicopter is able to land along the direct route of the land ambulance; and
 - ii. it would result in a significant reduction in transport time to the most appropriate hospital.
- 5. if the call's circumstances and patient(s) fail to meet the criteria set out in this standard and an air ambulance helicopter is known to be responding based on the merits of the initial request for ambulance service, contact the CACC/ACS and advise that an on-scene air ambulance helicopter response is not required and why it is not required.

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic.

Guideline

Airway / Breath.

Air Ambulance Helicopter Landing Site Safety and Coordination

Upon confirmation that the air ambulance helicopter is responding, the paramedic shall follow the guidelines set out by the Ornge Aviation Safety Department, which can be found on Ornge's "Aircraft Safety" website at: https://www.ornge.ca/aircraft-safety.

Other Use of Air Ambulance Helicopter

Cardiac/ Circula.

- Air ambulance helicopters are not permitted to respond to night calls which require a landing at a site other than night licensed airports, helipads or night approved remote landing sites.
 - Air ambulance helicopters are not permitted to conduct search and rescue calls.
- In cases where a land ambulance can reach the patient(s) and an on-scene response by air ambulance helicopter is appropriate, the ACO will assign a land ambulance and continue the land response until the flight crew requests that the land ambulance be cancelled.
- In cases where a land ambulance arrives on-scene prior to the air ambulance helicopter, paramedics shall inform the CACC/ACS as clinical events occur.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer

Medic.

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Intro

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Cardiac/ Circula.

LOC/ Pain/ Nausea

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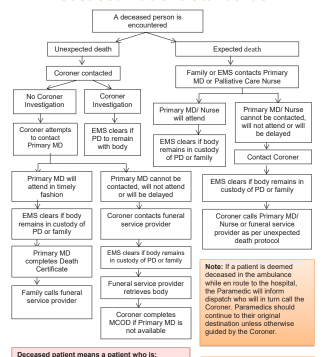
Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Deceased Patient Standards



Deceased patient means a patient who is:

- Obviously dead code 5
- ii) Subject to a MCOD presented to the paramedic
- iii) VSA and subject to a valid DNR
- iv) VSA and is subject to a Termination of Resuscitation Order
- v) VSA and is subject to a Withhold Resuscitation Order

Note: When a Termination of resuscitation Order is received, and the deceased person has not been removed from the place of death, paramedics should not remove the body, but rather they should follow the appropriate procedure as outlined.

Paramedic Prompt Card for Acute Stroke Protocol

This prompt card provides a quick reference of the Acute Stroke Protocol contained in the Basic Life Support Patient Care Standards (BLS PCS). Please refer to the BLS PCS for the full protocol.

Indications under the Acute Stroke Protocol

Redirect or transport to the closest or most appropriate Designated Stroke Centre* will be considered for patients who meet ALL of the following:

- 1. Present with a new onset of at least one of the following symptoms suggestive of the onset of an acute
 - Unilateral arm/leg weakness or drift.
 - Slurred speech or inappropriate words or mute.
 - Unilateral facial droop.
- 2. Can be transported to arrive at a Designated Stroke Centre within 6 hours of a clearly determined time of symptom onset or the time the patient was last seen in a usual state of health.
- 3. Perform a secondary screen for a Large Vessel Occlusion (LVO) stroke using the Los Angeles Motor Scale (LAMS) and inform the CACC/ACS to aid in the determination of the most appropriate destination.

*A Designated Stroke Center is a Regional Stroke Centre, District Stroke Centre or a Telestroke Centre regardless of EVT capability.

Contraindications under the Acute Stroke Protocol

ANY of the following exclude a patient from being transported under the Acute Stroke Protocol:

- 1. CTAS Level 1 and/or uncorrected airway, breathing or circulatory problem.
- Symptoms of the stroke resolved prior to paramedic arrival or assessment**.
- Blood sugar <3 mmol/L***.
- 4. Seizure at onset of symptoms or observed by paramedics.
- 5. Glasgow Coma Scale <10.
- 6. Terminally ill or palliative care patient.
- 7. Duration of out of hospital transport will exceed two hours.

**Patients whose symptoms improve significantly or resolve during transport will continue to be transported to a Designated Stroke Centre.

*** If symptoms persist after correction of blood glucose level, the patient is not contraindicated.

CACC/ACS will authorize the transport once notified of the patient's need for redirect or transport under the Acute Stroke Protocol.

Ontario 📆

Airway / Breath

Intro

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced

Research / Sp. Proj

Medical Refer.

Medic. Info

Cardiac/ Circula.

IOC/ Pain/ Nausea

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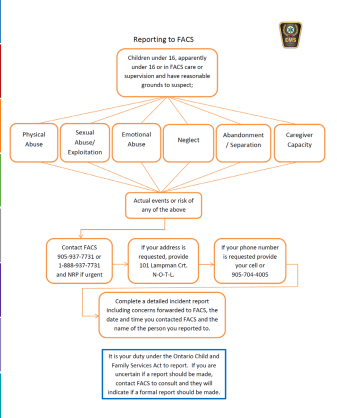
Research / Sp. Proj

Medical Refer

Medic Info.

Contact

Reporting to FACS Niagara



Paramedic Prompt Card for Sepsis



Paramedic Prompt Card for Sepsis Reference	YES	NO
Suspected or Confirmed Signs and Symptoms of Infection?		
Skin: Cellulitis, Wound, Burns		
Immunocompromised Neuro: LOC changes, Weakness, Indwelling Medical Device		
Chest: Cough, SOB, Recent Surgery/Invasive Procedure		
Abdomen: Pain, Vomiting, Diarrhea, History of Fever or Rigors (shakes)		
Urine: Dysuria, Frequency, Odour		
Age : ≥ 18		
At Least 2 OR MORE:		
Temperature: < 36° C OR ≥ 38° C		
Pulse: ≥ 90 bpm		
Respiratory Rate: ≥ 20bpm		
And at least ONE of the following		
Signs of Hypoperfusion (O2 Sat <92%)		
Systolic BP <90mmHg		
New Altered mental status		
Suggested Treatment		
IV access obtained		
Intravenous & Fluid Therapy Directive (bolus)		
Notify ED of *Sepsis Alert*		

Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Contact

Paramedic Prompt Card for Sepsis (NEMS)



Paramedic Prompt Card for Sepsis Reference	YES	NO	
Suspected or Confirmed Signs and Symptoms of			
Infection?			
Skin: Cellulitis, Wound, Burns			
Immunocompromised /Neuro: LOA changes,			
Weakness, Indwelling Medical Device , Chemotherapy			
Chest: Cough, SOB, Recent Surgery/Invasive			
Procedure			
Abdomen: Pain, Vomiting, Diarrhea with a history of			
fever or rigors			
 Urine: Dysuria, Frequency (increased or decreased), 			
Odour			
Age : ≥ 18			
At Least 2 OR MORE of the following:			
Temperature: < 36° C OR ≥ 38° C			
<i>Pulse</i> :≥ 90 bpm			
► Respiratory Rate:≥ 20bpm			
And at least ONE of the following			
 Signs of Hypoperfusion (mottled extremities, poor cap 			
refill, etc)			
Systolic BP <90mmHg			
New altered LOA			
If you answer yes to all of the above then Notify ED of *Sepsis Alert*			

Suggested Treatment

- IV access
- Intravenous & Fluid Therapy Directive
- If the patient clearly meets the Sepsis Alert AND they do not meet the Medical Directive for fluid therapy, consider contacting the BHP for IV fluid orders.

Airway /

Breath

Cardiac/

Circula.

IOC/

Pain/

Nausea

Proced

Research /

Sp. Proj

Medical Refer.

Medic. Info

Niagara EMS Hospital Destination Policy



Policy # IV 3.12a Hospital Destination Policy May 1, 2022

HOSPITAL DESTINATION POLICY - Niagara Region

The URGENT CARE CENTRE will only accept PATIENTS that meet the established auidelines

The Paramedic will:

Make a decision regarding receiving facility and transport the patient to that facility or an alternate facility as confirmed or directed by:

- > an ambulance dispatcher, or
- > an attending physician, with dispatch confirmation, or
- > a base hospital physician, with dispatch confirmation, or
- > approved local transfer guidelines, or
- the patient, with dispatch approval.

In the absence of direction, transport to the closest or most appropriate hospital emergency department capable of providing the medical care apparently required by the patient. The goal is to expedite time to definitive care. When there are two or more hospitals equal in time from the level 1 or 2 patient, the Paramedic may choose among available sites in consultation with NEMS Communications.

If in the paramedic's judgment, the patient can be managed en route the patient will be transported to the most appropriate hospital (as indicated below).

If the patient deteriorates during transport, and survival to the directed receiving facility is questionable, the paramedic will transport the patient to the closest or most appropriate hospital emergency department capable of providing the medical care immediately required by the patient. The paramedic will immediately notify dispatch of any destination change. and notify or ask dispatch to notify the initial and receiving facility.

Patient preference for a specific hospital, other than the closest, will be considered where resources permit based on clinical factors or continuity of care.

CONDITION	DESCRIPTION	DESTINATION
TRAUMA	Paramedics/ Dispatchers will consider the Air Ambulance Utilization Standard for FTT	Trauma Center/ Closest Emergency Department *
	All trauma patients meeting Field Trauma Triage (FTT) Standard Criteria where the incident location is within 60 minutes transport time to a Lead Trauma Centre will be transported to the Lead Trauma Centre in accordance with the guidelines (Policy IV-3.12h).	·
	*If transport time to Lead Trauma Centre will exceed 60 minutes, or survival to Lead Trauma Hospital is unlikely (see exception in Policy IV-	

-1-

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Policy # IV 3.12a Hospital Destination Policy May 1, 2022

ı	May 1, 2022		
		3.12h), patients meeting FTT criteria will be transported to the closest Emergency Department.	
	HEAD TRAUMA	All patients with head trauma & an altered LOC not meeting FTT Standard will be taken to the	Closest Emergency
İ	Hospitals with CT:	closest hospital with a functioning CT.	Department with a functioning CT
	GNG, SCS, WH Sites and WLMH in Niagara HGH Site in Hamilton	If they are in active resuscitation then the patient is to be transported to the closest ED.	(GNG, SCS, WH, WLMH and HGH)
	STROKE	Patients meeting the criteria of the Paramedic	Closest Stroke
l	EMERGENCIES	Prompt Card will be taken to the closest Stroke Centre for evaluation (attached)	Center
l	Stroke Centers: GNG Site and	Those stroke patients who do not meet the	
İ	Hamilton General Hospital	Paramedic Prompt Card criteria will be taken to the closest hospital with a functioning CT.	
	Hospitals with	If CT is down at the GNG Site, patients who meet the Provincial Paramedic Prompt Card criteria will	
l	GNG, SCG, WH Sites and WLMH in Niagara	be taken to the closest site with a functioning CT with "next on table" priority.	
	HGH in Hamilton	They will then be transported to the GNG Site for assessment by the Stroke Team (see attached Appendix A2-CT Downtime Contingency Plan for Stroke Thrombolysis (tPA).	
İ	SEXUAL ASSAULT	All victims of sexual assaults will go to the closest hospital for medical clearance.	Closest hospital for medical
l	ACCACE	·	clearance - then
		Following patient triage, registration, and physician assessment appropriate transfer arrangements to SCS/MGH will be made by the receiving site if the patient requires sexual assault services.	may require transfer to SCS or HGH as appropriate
	DIALYSIS	All hemo/ peritoneal dialysis with related	St. Catharines
	EMERGENCIES	complaints will be transported to SCS unless the patient is actively being resuscitated, patients will be transported to the closest hospital.	Site or St. Joseph's Health Care
		Consideration will be given to St. Joseph's Health Care Hamilton for patients picked up West of RR24	

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Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Policy # IV 3.12a Hospital Destination Policy May 1, 2022

OBSTETRICAL & GYNECOLOGICA L EMERGENCIES	Patients whose chief complaint is Obstetrical in nature will be taken to the SCS (or WLMH if closer) unless active resuscitation is in progress or in the case of a laboring patient a presenting fetal part is visible (e.g. crowning). These patients will be taken to the closest Emergency Department. If childbirth has occurred, and no active resuscitation is required, infant and mother should be transported to SCS or WLMH, whichever is closest. Note: WLMH should typically only be considered for patients greater than 36 weeks gestation. Patients whose presentation is highly suggestive of an ectopic pregnancy, for eg. sudden onset severe abdominal pain in a female of child bearing age, should also be considered for transport to SCS or WLMH if closer. Pregnant patients whose chief complaint is clearly NOT OB/GYN in nature will be transported under the appropriate destination for that complaint as outlined within this policy.	St. Catharines Site or WLMH, whichever is closest, unless active resuscitation in progress OR presenting fetal part is visible.	
ONCOLOGY and PALLIATIVE EMERGENCIES	Patients will go to the hospital where they have been receiving treatment within Niagara Region if they can be managed en route. Niagara's Regional Cancer Program is the SCS. (Consideration will be given to Juravinski in Hamilton for patients picked up West of RR24)	St. Catharines Site (consideration for Juravinski West of RR24)	

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Policy # IV 3.12a Hospital Destination Policy May 1, 2022

May 1, 2022				
PAEDIATRIC EMERGENCIES (less than 16 yrs. of age)	Paediatric patients triaged as Level 1 , or who require active resuscitation , will go to the closest hospital for immediate assessment and stabilization.	If active resuscitation go to closest hospital.		
	Non-complex Paediatric patients will be taken to the closest hospital or may be transported to a UCC in accordance with the Urgent Care Destination Criteria.	Complex patients go to St. Catharines Site or MUMC depending on location		
	Complex patients, such as those with indwelling medical devices, with medically complex histories or injuries, or who are currently receiving treatment at St. Catharines Site, should be transported to the closest hospital with a pediatrician available (SCS in Niagara, MUMC in Hamilton) if the patient can be managed during transport.			
	All other patients will be transported to the closest appropriate hospital as outlined in this policy (for example, orthopedics or trauma).			
MENTAL HEALTH EMERGENCIES	Patients of all ages where mental illness is the primary problem will be taken to a schedule 1 facility: SCS in Niagara, or St. Joseph's Healthcare in Hamilton if closer. Patients should be taken to the closest of the two sites.	If primary problem is medical go to closest hospital.		
	Consideration for previous treatment history with a facility may be considered in choosing an appropriate destination.	If Mental Illness is the primary problem then go to St. Catharines		
	Patients with a history of mental illness, but in whom the primary problem is medical (i.e. overdose etc.) or surgical emergency will go to the closest appropriate hospital as outlined elsewhere	Site, or SJHH if closer.		
	in this policy.			
ORTHOPEDIC EMERGENCIES	Patients with major orthopedic emergencies (i.e. long bone fracture, spinal or pelvic fracture, open fracture or gross deformity) will be taken to the closest appropriate hospital i.e. where there is an	Major: Closest hospital with Ortho (peds to SCS or MUMC)		
	Orthopedic Surgeon on-call if they can be managed en route. This includes HGH to the West. Patients under 16 should be transported to SCS (MUMC if closer)	Minor: Closest hospital or UCC		

Contact

- 4 -

Policy # IV 3.12a Hospital Destination Policy

Policy # IV 3.12a Hospital Destination Policy

May 1, 2022

May 1, 2022

Patients with minor orthopedic emergencies (i.e. isolated orthopedic injury, fractured wrist, ankle etc.) will be taken to the closest hospital ED or	
UCC if they meet the Urgent Care Centre Destination Criteria.	

Revised: May 1, 2022

Airway / Breath

Cardiac/ Circula.

IOC/ Pain/ Nausea

PARAMEDIC PROMPT CARD Niagara Regional Acute Stroke Protocol

Refer to current Paramedic Prompt Card for Acute Stroke Protocol contained within the current Basic Life Support Patient Care Standards.

The closest Stroke Centre is defined in the CAD.

Notify the Receiving Hospital that they will be receiving a "Stroke Alert" patient that meets the Acute Stroke Protocol.

Transport CTAS Level 2 to the Emergency Department of the closest Stroke Centre.

Proced

Research / Sp. Proj

Medical Refer.

Medic. Info

Cardiac/ Circula.

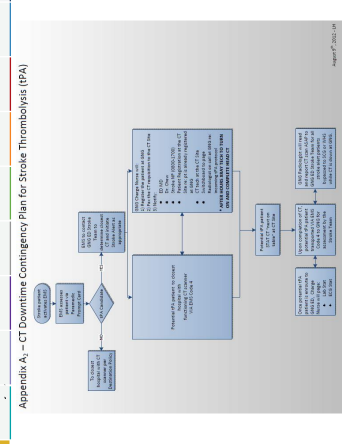
LOC/ Pain/ Nausea

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Research / Sp. Proj

Medical Refer.

Medic. Info.





"The Canadian CSPINE Rule"

Airway / Breath

> Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic.

Contact

Inclusion Criteria 1. Any ONE High-Risk factor which Alert (GCS 15) mandates immobilization? Stable (SBP>90, RR 10-24) · Age ≥65 years Acute Blunt Injury (<48 hrs) · Dangerous Mechanism* · Numbness or tingling in extremities **L** No. 2. Any ONE Low-Risk factor which allows safe assessment of range of motion? • Rear-ended in Simple Rear-end MVC** USE C-SPINE No · Ambulatory at any time at scene **IMMOBILIZATION** No neck pain at scene when asked (answer "yes" if no pain) · No pain during midline c-spine palpation (answer "yes" if no pain) Yes 3. Patient voluntarily able to Actively **Exclusion Criteria** Rotate neck 45° left and right when Boarded & Collared for Other Reasons requested, regardless of pain? Aae<8 Penetrating Trauma Yes Acute Paralysis Known Vertebral Disease NO C-SPINE IMMOBILIZATION Referred from another Hospital *Dangerous Mechanism: **Simple Rearend MVC Excludes: • fall from elevation ≥3 feet/5 stairs · pushed into oncoming traffic · axial load to head, e.g. diving hit by bus/large truck

rollover

hit by high speed vehicle (≥100km/h)

MVC: rollover, ejection, high speed (≥100km/h)

· bicycle collision with object, e.g. post, car

· motorized recreational vehicles, e.g. ATV, snowmobile

Airway / Breath.

Cardiac/

Circula.

IOC/

Pain/

Nausea

Proced.

Research/

Sp. Proj

Medical

Refer

Medic. Info.

STEMI Hospital Bypass Prompt Card

This prompt card provides a quick reference of the STEMI Hospital Bypass Protocol contained in the Basic Life Support Patient Care Standards (BLS PCS). Please refer to the BLS PCS for the full protocol.

Indications under the STEMI Hospital Bypass Protocol

Transport to a PCI centre will be considered for patients who meet ALL of the following:

- ≥18 years of age.
- 2. Chest pain or equivalent consistent with cardiac ischemia/myocardial infarction.
- 3. Time from onset of current episode of pain <12 hours.
- 4. 12-lead ECG indicates an acute AMI/STEMI*:
 - a. At least 2 mm ST-elevation in leads V1-V3 in at least two contiguous leads; AND/OR
 - b. At least 1 mm ST-elevation in at least two other anatomically contiguous leads; OR
 - c. 12-lead ECG computer interpretation of STEMI and paramedic agrees.

*Once activated, continue to follow the STEMI Hospital Bypass Protocol even if the ECG normalizes.

Contraindications under the STEMI Hospital Bypass Protocol

ANY of the following exclude a patient from being transported under the STEMI Hospital Bypass Protocol:

- 1. CTAS 1 and the paramedic is unable to secure patient's airway or ventilate.
- 2. 12-lead ECG is consistent with a LBBB, ventricular paced rhythm, or any other STEMI imitator
- Transport to a PCI centre ≥60 minutes from patient contact.**
- 4. Patient is experiencing a complication requiring PCP diversion:**
 - Moderate to severe respiratory distress or use of CPAP.
 - b. Hemodynamic instability or symptomatic SBP <90 mmHg at any point.
 - c. VSA without ROSC.
- 5. Patient is experiencing a complication requiring ACP diversion:**
 - Ventilation inadequate despite assistance.
 - b. Hemodynamic instability unresponsive/not amenable to ACP treatment/management.
 - c. VSA without ROSC.

CACC/ACS will authorize the transport once notified of the patient's need for bypass under the STEMI Hospital Bypass Protocol.

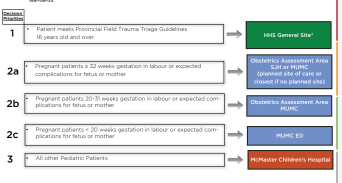
Ontario 😿

^{**}The interventional cardiology program may still permit the transport to the PCI centre.

Pediatric Patient Priority System (PPS)



Pediatric patients (less than 18 years) will be transported according to the Basic Life Support Patient Care Standards, Patient Transport Standard. The following presentations should be taken to the facility listed as the most appropriate hospital capable of providing the medical care apparently required by the patient. VSA, pre-arrest or unresolved airway compromise patients should be transported to the closest facility unless otherwise directed by provincial guidelines/ standards.



Suspected Ebola Virus Disease (EVD) disease patients must be considered according to the tool attached *In any case that a regional hospital is closed to any incoming patients (i.e. fire in the hospital), CACC will decide the hospital destination.

Airway / Breath.

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

IOC/ Pain/

Adult Patient Priority System (PPS) (HPS) Adult patients 18 years and older will be transported according to the Basic Life Support Patient Care Standards, Patient Transport Airway / Standard. The following presentations should be taken to the facility listed as the most appropriate hospital capable of providing the medical care apparently required by the patient. VSA, pre-arrest or unresolved airway compromise patients should be Breath. transported to the closest facility unless otherwise directed by provincial guidelines/standards. Patient meets Field Trauma Triage Guidelines, including pregnant patient meeting Field Trauma Triage Guidelines Possible ST Elevation MI (Provincial Paramedic Prompt Card) Acute Stroke (Provincial Paramedic Prompt Card) HHS General Site* Major Burn >25% Total Body Surface or airway problems Smoke Inhalation Injury with altered LOC Cardiac/ Diving/Decompression Incidents Circula. Dialysis patient St. Joseph's Healthcare Psychiatric emergency (as per Recognition Tool) SJH or MUMC (planned site of care or closest if no planned site) Pregnant patients ≥ 32 weeks gestation in labour or expected 3a complications for fetus or mother Nausea Pregnant patients 20-31 weeks destation in labour or expected 3b complications for fetus or mother Proced. Pregnant patients < 20 weeks destation in labour or expected complications for fetus or mother 3c All other Pregnant patients regardless of gestational age with non-FTTG injury or other medical concern HHS General Site or Known or suspected Sexual Assault HHS Juravinski Site Research/ Sp. Proj Possible GI Bleed (as per Recognition Tool) St. Joseph's Healthcare* Possible Hip Fracture (as per Recognition Tool) or HHS Juravinski Site* St. Joseph's Healthcare* 6 (as per Recognition Tool) or HHS General Site* Medical Refer UCC Patients (St. Joseph's King Street East Campus UCC, and HHS Any "arranged" ED or direct Main Street West UCC) transported to the "arranged" Emergency 7 to any "arranged" unit (with Department for continuation of the patient care. immediate transfer of care). Patients with a recent history at a particular hospital for a related Facility with most Medic problem (defined as inpatient within 14 days) recent history (as defined). Any "arranged" ED or direct Attending physician has made arrangements, as confirmed by Hamilton CACC with the receiving hospital and the "accepting" to any "arranged" hospital unit. physician identified.

NOTE: For Decision Priorities #7 through #9, CACC will endeavor to distribute patients in a manner that facilitates equity and prompt transfer Suspected Ebola Virus Disease (EVD) disease patients must be considered according to the tool attached

As directed by CACC

considering all factors

"In any case that a regional hospital is closed to any incoming patients (i.e. fire in the hospital). CACC will decide the hospital destination.

Contact

Info.

10

All other patients.

GI Bleed Recognition Tool (HPS)



For the purposes of the Patient Priority System:

Patients with possible "GI bleeds" (gastrointestinal bleeding) recognized by the guidelines below should be transported to the appropriate Emergency Department (St. Joseph's Healthcare or HHS Juravinski Site) as directed by CACC.

INCLUSION

The patient must be; ≥ 18 years of age and meet the following:

- 1. Vomiting blood (hematemesis) bright red blood. dark red blood, dark brown/black blood ("coffee grounds") or blood clots.
- 2. Passing red blood rectally (hematochezia) bright red blood, dark red blood or blood clots (with or without stools)
- 3. Passing black stools (melena) sticky, black, "tarry", stools with a typical foul smell - may be mixed with red or maroon blood.

EXCLUSION

Patients < 18 years should be transported as per the Pediatric Destination Determination Guidelines and not according to this Tool.

Airway / Breath

Intro

Cardiac/ Circula.

IOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info

Contact

Education notes:

Relevant history:

If a patient with a possible "GI bleed" has an extensive history with one site (eg: such as post operative, oncology, dialysis, multiple admissions, or discharged patient), it would be preferable for the patient to be transported to that site (excluding McMaster Children's Hospital or HHS Hamilton General Site).

Airway / Breath.

Isolated Hip Fracture Recognition Tool (HPS)



For the purposes of the Patient Priority System:

Patients with possible "isolated" hip fracture recognized by the guidelines below should be transported to the Emergency Department as directed by CACC (St. Joseph's Healthcare or HHS Juravinski Site).

Cardiac/ Circula.

IOC/

Pain/

Nausea

INCLUSION

Mechanism: Fall from sitting (chair), bed, or standing (not height or MVC); may have other minor injuries (i.e. contusions); AND

History of: Pain in hip or groin at rest or with patient initiated movement (paramedic should not intentionally move joint); AND

Examination: May have externally rotated and/or shortened leg.

EXCLUSION

- 1. Patient meets the Trauma Triage Guidelines
- Patient with hip joint replacement on same side (Pt should be transported to site of original joint replacement surgery. If original site is unknown normal distribution guidelines will apply).

Proced.

Research/ Sp. Proj

Education notes:

1. "Isolated" hip fracture: Refers to no other recognized significant injuries.

Mechanisr

The intention of the above listed mechanism is to select those patients that are unlikely to have additional injuries (significant trauma mechanism). Although the tool states fall from sitting, lying, standing, this may also include a single step or curb but is meant to exclude more significant falls.

Relevant history:

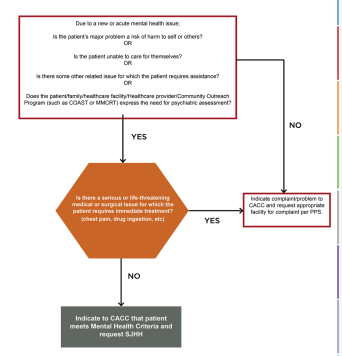
If a patient with a possible hip fracture has an extensive history with one site (i.e. such as post-operative, oncology, dialysis, multiple admissions, or discharged patient), it would be preferable for the patient to be transported to that site (excluding McMaster Children's Hospital or HHS Hamilton General Site).

Medical Refer.

Medic. Info.

Psychiatric Emergency Recognition Tool (HPS)





Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

Research/ Sp. Proj

Medical Refer.

Medic. Info.

Airway / Breath.

Cardiac/ Circula.

IOC/

Pain/

Nausea

Musculoskeletal Injury Recognition Tool (HPS)



For the purposes of Patient Priority System:

Patients with suspected significant orthopedic fractures which might require immediate surgery (excluding hip) by the guidelines below should be transported to the Emergency Departments of St. Joseph's Hospital or Hamilton General Hospital as directed by CACC.

INCLUSION

Adult patients (≥18) with:

- 1. Suspected "open" fracture of any limb, OR
- 2. Severe bony deformity of an injured lower limb

EXCLUSION

- 1. Patient's injury is at site of known joint replacement (prosthetic joint), then transport to the Emergency Department to the site where the joint replacement surgery was performed or the Juravinski or St. Joseph's Hospital as directed by CACC.
- 2. Receiving active oncology treatment at the Juravinski Cancer Clinic, transport to the Juravinski Emergency Department.

Research/

Sp. Proj

Medical

Refer

Medic

Info.

Proced.

Education notes:

- 1, If Patient meets the Provincial Trauma Triage Guidelines, then transport to Hamilton General Hospital as directed by CACC.
 - 2. If Patient meets the Possible Hip Fracture Identification Tool, preferentially follow that tool, then transport to the Emergency Department of the Juravinski or St. Joseph's Hospital as directed by CACC.
- 3. "Open" fracture or compound fracture: Refers to a fracture with an associated wound. This can include circumstances where the bone fragments can be seen protruding through a wound, where there is a large skin defect or even just a small puncture sized wound where the bone may have penetrated the skin but is no longer visible. Any open injury (other than an abrasion) associated with a suspected fracture can be considered a suspected "open" fracture for the purposes of this auideline.
- 4. The Juravinski Hospital will continue to treat pathological fractures associated with a malignancy
- 5. All Sites, including the Juravinski Hospital, will continue to manage patients with fractures not requiring immediate surgery, dislocations and soft tissue injuries.

Ebola Virus Disease (EVD) Screening Recognition Tool



For the purposes of the Patient Priority System:

Patients who are screened as positive (suspected EVD) using the most current Ministry of Health and Long Term Care (MOHLTC) EVD Screening Tool, and who meet specific destination protocol criteria, will be preferentially transported as indicated below:

Adult patient ≥18 years of age and screened positive for EVD:

. For Decision Priority 1 through 4, follow the current Adult PPS by transporting the patient to the identified destination as per normal practice.

For Decision Priority 5 through 10, transport the adult patient to the Juravinski Hospital

Pediatric patient <18 years of age and screened positive for EVD:

• For all Decision Priority criteria follow the current Pediatric PPS by transporting the patient to the identified destination as per normal practice.

Education Notes:

- 1. When a patient has screened positive for EVD, a patch to notify the receiving facility must be completed by the Paramedics regardless of transport priority.
- 2. The following hospitals are designated EVD testing sites although the ambulance destination decision will follow the direction above:
 - Juravinski Hospital Adult patients (≥18 years of age)
 - McMaster Children's Hospital Pediatric patients (<18 years of age)

Airway /

Intro

Cardiac/ Circula.

Breath

IOC/ Pain/ Nausea

Proced

Research / Sp. Proj

Medical Refer.

Medic. Info

Airway / Breath

Radio Channel Change Locations



Hamilton

QEW and Fifty Road=====NIA REG2 COM, contact Hamilton CACC

London

QEW and Fifty Road=====NIA REG2 COM, contact Hamilton CACC

Hwy 403 and County Road 25 (Middle Townline Road)=====NIA MOH ZN 1, contact London CACC

This is about 15-20 km west of Brantford

Mississauga

QEW and Fifty Road====NIA REG2 COM, contact Hamilton CACC

QEW and Hwy 403 (base of Burlington Skyway)=====NIA MOH ZN 1, contact Mississauga CACC

Toronto

QEW and Fifty Road====NIA REG2 COM, contact Hamilton CACC

QEW and Hwy 403 (base of Burlington Skyway)=====NIA MOH ZN 1, contact Mississauga CACC

QEW and Hwy 427====NIA PROV COM, contact Toronto CACC

When returning, the locations for changing back are the same.

If transporting a patient on return to Niagara, switch to NIA TAC 1 at Fifty Road. If you are returning empty, switch to NIA North at Fifty Road.

All channels are within the NIA folder and can be found by simply turning the Channel Selector.

> Opt Zone NIA REG2 COM

Cardiac/

Circula.

IOC/ Pain/ Nausea

Proced.

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Medic Info.











www.sepsis-prealert.ca



Airway / Breath.

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Contact

FAST Sepsis Pre-Alert for GWPS, HPS, and ROWPS

Do you suspect or know there is an infection? If yes, apply ParaHEWS (below) If ParaHEWS ≥5: notify receiving hospital of "Sepsis Pre-Alert" and Apply Capnography

Physiological Parameters	8	2	1	0	1	2	က
Heart Rate / Pulse		<41	41-50	51-100	101-110	111-130	>131
Systolic BP	<71	71-90		91-170		171-200	>201
Respiratory Rate	8 >	8-13		14-20		21-30	≥31
Temperature (C)	<35		35.0-36.0	36.1-37.9 (or not available)	38.0-39.0	≥39.1	
O ₂ Saturation	<85		85-92	≥93			
O ₂ Therapy				Room Air	O ₂ via nasal prongs		O ₂ via face mask
Change in CNS from Baseline		New Confusion		Alert or Usual Self	Voice	Pain	Not responsive

STEMI Protocol Pearls



Airway / Breath.

Cardiac/

LOC/ Pain/ Nausea

Proced.

Research / Sp. Proj

Medical Refer.

Medic.

Contact

Sympt oms

PAIN

Pain can be typical or atypical (but not only non-specific symptoms of dyspnea, nausea, fatique, etc)

ACUTE

An acute history of symptoms of < 12 hours



ECG

QUALITY

- Ensure good quality ECG
- · Shave chest
- No moving/talking

REPEAT

If negative, do serial ECGs

(1) before treatment

(2) in ambulance prior to leaving scene
(3) in ambulance prior to moving into ED

CAUTION

ECGs can be tricky, rule out mimics If not certain, go to closest appropriate ED







Geography

60 MINUTES

Maximum 60 minutes from first medical contact to PCI centre

If you are quicker on scene (eg: 15 minutes), this will allow longer transport time (eg: 45 minutes)



BOUNDARIES

Know the PCI centres in your area CACC may be able to assist

1-844-832-6830

Brampton 1-416-747-3500,1

St. Mary's 1-519-653-4074 Southlake 1-905-952-2466

Trillium 1-888-493-3568

Pr epar e

CALITION

Caution with nitro and morphine

Neither of these medications are life-saving in STEMI patients & can cause adverse events

"PADS ON"

Defibrillation pads are placed on all patients with suspected STEMI



BE READY

Be familiar with the common complications

- · dysrhythmias
- · pump failure
- cardiac arrest

Be ready to manage them



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Airway / Breath.

Cardiac/ Circula.

LOC/ Pain/ Nausea

Proced.

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Medic. Info.

LIST OF MANDATORY PROVINCIAL PATCH POINTS

Medical Cardiac Arrest

• TOR

Trauma Cardiac Arrest

• Trauma TOR

IV & Fluid Therapy

• Fluid bolus for hypotensive patients <12 years of age with suspected DKA

LIST OF MANDATORY LOCAL PATCH POINTS

- Special Project Palliative Care Medical Directives
- Research Project Palliative Care Medical Directives





Medication Safety Starts with You

When you see the "5Rs" symbol throughout this guidebook, it is a reminder to always confirm:

- **ORIGHT PATIENT**
 - **O** RIGHT **DRUG**
 - **O** RIGHT **DOSE**
 - **O** RIGHT **ROUTE**
 - **O** RIGHT **TIME**

